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NOVEL NUCLEIC ACIDS AND POLYPEPTIDES

1. TECHNICAL FIELD

The present invention provides novel polynucleotides and proteins encoded by such polynucleotides, along with uses for these polynucleotides and proteins, for example in therapeutic, diagnostic and research methods.

2. BACKGROUND

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Technology aimed at the discovery of protein factors (including e.g., cytokines, such as lymphokines, interferons, circulating soluble factors, chemokines, and interleukins) has matured rapidly over the past decade. The now routine hybridization cloning and expression cloning techniques clone novel polynucleotides "directly" in the sense that they rely on information directly related to the discovered protein (i.e., partial DNA/amino acid sequence of the protein in the case of hybridization cloning; activity of the protein in the case of expression cloning). More recent "indirect" cloning techniques such as signal sequence cloning, which isolates DNA sequences based on the presence of a now well-recognized secretory leader sequence motif, as well as various PCR-based or low stringency hybridization-based cloning techniques, have advanced the state of the art by making available large numbers of DNA/amino acid sequences for proteins that are known to have biological activity, for example, by virtue of their secreted nature in the case of leader sequence cloning, by virtue of their cell or tissue source in the case of PCR-based techniques, or by virtue of structural similarity to other genes of known biological activity.

Identified polynucleotide and polypeptide sequences have numerous applications in, for example, diagnostics, forensics, gene mapping; identification of mutations responsible for genetic disorders or other traits, to assess biodiversity, and to produce many other types of data and products dependent on DNA and amino acid sequences.

3. SUMMARY OF THE INVENTION

The compositions of the present invention include novel isolated polypeptides, novel isolated polynucleotides encoding such polypeptides, including recombinant DNA molecules, cloned genes or degenerate variants thereof, especially naturally occurring variants such as allelic variants, antisense polynucleotide molecules, and antibodies that specifically recognize

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one or more epitopes present on such polypeptides, as well as hybridomas producing such antibodies.

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The compositions of the present invention additionally include vectors, including expression vectors, containing the polynucleotides of the invention, cells genetically engineered to contain such polynucleotides and cells genetically engineered to express such polynucleotides.

The present invention relates to a collection or library of at least one novel nucleic acid sequence assembled from expressed sequence tags (ESTs) isolated mainly by sequencing by hybridization (SBH), and in some cases, sequences obtained from one or more public databases. The invention relates also to the proteins encoded by such polynucleotides, along with therapeutic, diagnostic and research utilities for these polynucleotides and proteins. These nucleic acid sequences are designated as SEQ ID NO: 1-341. The polypeptides sequences are designated SEQ ID NO: 342-682. The nucleic acids and polypeptides are provided in the Sequence Listing. In the nucleic acids provided in the Sequence Listing, A is adenosine; C is cytosine; G is guanine; T is thymine; and N is unknown or any of the four bases.

The nucleic acid sequences of the present invention also include, nucleic acid sequences that hybridize to the complement of SEQ ID NO: 1-341 under stringent hybridization conditions; nucleic acid sequences which are allelic variants or species homologues of any of the nucleic acid sequences recited above, or nucleic acid sequences that encode a peptide comprising a specific domain or truncation of the peptides encoded by SEQ ID NO: 1-341. A polynucleotide comprising a nucleotide sequence having at least 90% identity to an identifying sequence of SEQ ID NO: 1-341 or a degenerate variant or fragment thereof. The identifying sequence can be 100 base pairs in length.

The nucleic acid sequences of the present invention also include the sequence information from the nucleic acid sequences of SEQ ID NO: 1-341. The sequence information can be a segment of any one of SEQ ID NO: 1-341 that uniquely identifies or represents the sequence information of SEQ ID NO: 1-341.

A collection as used in this application can be a collection of only one polynucleotide. The collection of sequence information or identifying information of each sequence can be provided on a nucleic acid array. In one embodiment, segments of sequence information are provided on a nucleic acid array to detect the polynucleotide that contains the segment. The array can be designed to detect full-match or mismatch to the polynucleotide that contains the segment. The collection can also be provided in a computer-readable format.

This invention also includes the reverse or direct complement of any of the nucleic acid sequences recited above; cloning or expression vectors containing the nucleic acid sequences; and host cells or organisms transformed with these expression vectors. Nucleic acid sequences (or their reverse or direct complements) according to the invention have numerous applications in a variety of techniques known to those skilled in the art of molecular biology, such as use as hybridization probes, use as primers for PCR, use in an array, use in computer-readable media, use in sequencing full-length genes, use for chromosome and gene mapping, use in the recombinant production of protein, and use in the generation of anti-sense DNA or RNA, their chemical analogs and the like.

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In a preferred embodiment, the nucleic acid sequences of SEQ ID NO: 1-341 or novel segments or parts of the nucleic acids of the invention are used as primers in expression assays that are well known in the art. In a particularly preferred embodiment, the nucleic acid sequences of SEQ ID NO: 1-341 or novel segments or parts of the nucleic acids provided herein are used in diagnostics for identifying expressed genes or, as well known in the art and exemplified by Vollrath et al., Science 258:52-59 (1992), as expressed sequence tags for physical mapping of the human genome.

The isolated polynucleotides of the invention include, but are not limited to, a polynucleotide comprising any one of the nucleotide sequences set forth in SEQ ID NO: 1-341; a polynucleotide comprising any of the full length protein coding sequences of SEQ ID NO: 1-341; and a polynucleotide comprising any of the nucleotide sequences of the mature protein coding sequences of SEQ ID NO: 1-341. The polynucleotides of the present invention also include, but are not limited to, a polynucleotide that hybridizes under stringent hybridization conditions to (a) the complement of any one of the nucleotide sequences set forth in SEQ ID NO: 1-341; (b) a nucleotide sequence encoding any one of the amino acid sequences set forth in the Sequence Listing; (c) a polynucleotide which is an allelic variant of any polynucleotides recited above; (d) a polynucleotide which encodes a species homolog (e.g. orthologs) of any of the proteins recited above; or (e) a polynucleotide that encodes a polypeptide comprising a specific domain or truncation of any of the polypeptides comprising an amino acid sequence set forth in the Sequence Listing.

The isolated polypeptides of the invention include, but are not limited to, a polypeptide comprising any of the amino acid sequences set forth in SEQ ID NO: 342-682; or the corresponding full length or mature protein. Polypeptides of the invention also include polypeptides with biological activity that are encoded by (a) any of the polynucleotides having a

nucleotide sequence set forth in SEQ ID NO: 1-341; or (b) polynucleotides that hybridize to the complement of the polynucleotides of (a) under stringent hybridization conditions. Biologically or immunologically active variants of any of the polypeptide sequences in the Sequence Listing, and "substantial equivalents" thereof (e.g., with at least about 65%, 70%, 75%, 80%, 85%, 90%, 95%, 98% or 99% amino acid sequence identity) that preferably retain biological activity are also contemplated. The polypeptides of the invention may be wholly or partially chemically synthesized but are preferably produced by recombinant means using the genetically engineered cells (e.g. host cells) of the invention.

The invention also provides compositions comprising a polypeptide of the invention. Polypeptide compositions of the invention may further comprise an acceptable carrier, such as a hydrophilic, e.g., pharmaceutically acceptable, carrier.

The invention also provides host cells transformed or transfected with a polynucleotide of the invention.

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The invention also relates to methods for producing a polypeptide of the invention comprising growing a culture of the host cells of the invention in a suitable culture medium under conditions permitting expression of the desired polypeptide, and purifying the polypeptide from the culture or from the host cells. Preferred embodiments include those in which the protein produced by such process is a mature form of the protein.

Polynucleotides according to the invention have numerous applications in a variety of techniques known to those skilled in the art of molecular biology. These techniques include use as hybridization probes, use as oligomers, or primers, for PCR, use for chromosome and gene mapping, use in the recombinant production of protein, and use in generation of anti-sense DNA or RNA, their chemical analogs and the like. For example, when the expression of an mRNA is largely restricted to a particular cell or tissue type, polynucleotides of the invention can be used as hybridization probes to detect the presence of the particular cell or tissue mRNA in a sample using, e.g., in situ hybridization.

In other exemplary embodiments, the polynucleotides are used in diagnostics as expressed sequence tags for identifying expressed genes or, as well known in the art and exemplified by Vollrath et al., Science 258:52-59 (1992), as expressed sequence tags for physical mapping of the human genome.

The polypeptides according to the invention can be used in a variety of conventional procedures and methods that are currently applied to other proteins. For example, a polypeptide of the invention can be used to generate an antibody that specifically binds the

polypeptide. Such antibodies, particularly monoclonal antibodies, are useful for detecting or quantitating the polypeptide in tissue. The polypeptides of the invention can also be used as molecular weight markers, and as a food supplement.

Methods are also provided for preventing, treating, or ameliorating a medical condition which comprises the step of administering to a mammalian subject a therapeutically effective amount of a composition comprising a polypeptide of the present invention and a pharmaceutically acceptable carrier.

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In particular, the polypeptides and polynucleotides of the invention can be utilized, for example, in methods for the prevention and/or treatment of disorders involving aberrant protein expression or biological activity.

The present invention further relates to methods for detecting the presence of the polynucleotides or polypeptides of the invention in a sample. Such methods can, for example, be utilized as part of prognostic and diagnostic evaluation of disorders as recited herein and for the identification of subjects exhibiting a predisposition to such conditions. The invention provides a method for detecting the polynucleotides of the invention in a sample, comprising contacting the sample with a compound that binds to and forms a complex with the polynucleotide of interest for a period sufficient to form the complex and under conditions sufficient to form a complex and detecting the complex such that if a complex is detected, the polynucleotide of interest is detected. The invention also provides a method for detecting the polypeptides of the invention in a sample comprising contacting the sample with a compound that binds to and forms a complex with the polypeptide under conditions and for a period sufficient to form the complex and detecting the formation of the complex such that if a complex is formed, the polypeptide is detected.

The invention also provides kits comprising polynucleotide probes and/or monoclonal antibodies, and optionally quantitative standards, for carrying out methods of the invention. Furthermore, the invention provides methods for evaluating the efficacy of drugs, and monitoring the progress of patients, involved in clinical trials for the treatment of disorders as recited above.

The invention also provides methods for the identification of compounds that modulate (i.e., increase or decrease) the expression or activity of the polynucleotides and/or polypeptides of the invention. Such methods can be utilized, for example, for the identification of compounds that can ameliorate symptoms of disorders as recited herein. Such methods can include, but are not limited to, assays for identifying compounds and other

substances that interact with (e.g., bind to) the polypeptides of the invention. The invention provides a method for identifying a compound that binds to the polypeptides of the invention comprising contacting the compound with a polypeptide of the invention in a cell for a time sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a reporter gene sequence in the cell; and detecting the complex by detecting the reporter gene sequence expression such that if expression of the reporter gene is detected the compound the binds to a polypeptide of the invention is identified.

The methods of the invention also provide methods for treatment which involve the administration of the polynucleotides or polypeptides of the invention to individuals exhibiting symptoms or tendencies. In addition, the invention encompasses methods for treating diseases or disorders as recited herein comprising administering compounds and other substances that modulate the overall activity of the target gene products. Compounds and other substances can effect such modulation either on the level of target gene/protein expression or target protein activity.

The polypeptides of the present invention and the polynucleotides encoding them are also useful for the same functions known to one of skill in the art as the polypeptides and polynucleotides to which they have homology (set forth in Table 2); for which they have a signature region (as set forth in Table 3); or for which they have homology to a gene family (as set forth in Table 4). If no homology is set forth for a sequence, then the polypeptides and polynucleotides of the present invention are useful for a variety of applications, as described herein, including use in arrays for detection.

4. DETAILED DESCRIPTION OF THE INVENTION

25 4.1 **DEFINITIONS**

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It must be noted that as used herein and in the appended claims, the singular forms "a", "an" and "the" include plural references unless the context clearly dictates otherwise.

The term "active" refers to those forms of the polypeptide which retain the biologic and/or immunologic activities of any naturally occurring polypeptide. According to the invention, the terms "biologically active" or "biological activity" refer to a protein or peptide having structural, regulatory or biochemical functions of a naturally occurring molecule. Likewise "immunologically active" or "immunological activity" refers to the capability of the

natural, recombinant or synthetic polypeptide to induce a specific immune response in appropriate animals or cells and to bind with specific antibodies.

The term "activated cells" as used in this application are those cells which are engaged in extracellular or intracellular membrane trafficking, including the export of secretory or enzymatic molecules as part of a normal or disease process.

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The terms "complementary" or "complementarity" refer to the natural binding of polynucleotides by base pairing. For example, the sequence 5'-AGT-3' binds to the complementary sequence 3'-TCA-5'. Complementarity between two single-stranded molecules may be "partial" such that only some of the nucleic acids bind or it may be "complete" such that total complementarity exists between the single stranded molecules. The degree of complementarity between the nucleic acid strands has significant effects on the efficiency and strength of the hybridization between the nucleic acid strands.

The term "embryonic stem cells (ES)" refers to a cell that can give rise to many differentiated cell types in an embryo or an adult, including the germ cells. The term "germ line stem cells (GSCs)" refers to stem cells derived from primordial stem cells that provide a steady and continuous source of germ cells for the production of gametes. The term "primordial germ cells (PGCs)" refers to a small population of cells set aside from other cell lineages particularly from the yolk sac, mesenteries, or gonadal ridges during embryogenesis that have the potential to differentiate into germ cells and other cells. PGCs are the source from which GSCs and ES cells are derived. The PGCs, the GSCs and the ES cells are capable of self-renewal. Thus these cells not only populate the germ line and give rise to a plurality of terminally differentiated cells that comprise the adult specialized organs, but are able to regenerate themselves.

The term "expression modulating fragment," EMF, means a series of nucleotides which modulates the expression of an operably linked ORF or another EMF.

As used herein, a sequence is said to "modulate the expression of an operably linked sequence" when the expression of the sequence is altered by the presence of the EMF. EMFs include, but are not limited to, promoters, and promoter modulating sequences (inducible elements). One class of EMFs are nucleic acid fragments which induce the expression of an operably linked ORF in response to a specific regulatory factor or physiological event.

The terms "nucleotide sequence" or "nucleic acid" or "polynucleotide" or "oligonculeotide" are used interchangeably and refer to a heteropolymer of nucleotides or the sequence of these nucleotides. These phrases also refer to DNA or RNA of genomic or

synthetic origin which may be single-stranded or double-stranded and may represent the sense or the antisense strand, to peptide nucleic acid (PNA) or to any DNA-like or RNA-like material. In the sequences herein A is adenine, C is cytosine, T is thymine, G is guanine and N is A, C, G or T (U). It is contemplated that where the polynucleotide is RNA, the T (thymine) in the sequences provided herein is substituted with U (uracil). Generally, nucleic acid segments provided by this invention may be assembled from fragments of the genome and short oligonucleotide linkers, or from a series of oligonucleotides, or from individual nucleotides, to provide a synthetic nucleic acid which is capable of being expressed in a recombinant transcriptional unit comprising regulatory elements derived from a microbial or viral operon, or a eukaryotic gene.

The terms "oligonucleotide fragment" or a "polynucleotide fragment", "portion," or "segment" or "probe" or "primer" are used interchangeably and refer to a sequence of nucleotide residues which are at least about 5 nucleotides, more preferably at least about 7 nucleotides, more preferably at least about 11 nucleotides, more preferably at least about 11 nucleotides and most preferably at least about 17 nucleotides. The fragment is preferably less than about 500 nucleotides, preferably less than about 200 nucleotides, more preferably less than about 100 nucleotides, more preferably less than about 50 nucleotides and most preferably less than 30 nucleotides. Preferably the probe is from about 6 nucleotides to about 200 nucleotides, preferably from about 15 to about 50 nucleotides, more preferably from about 17 to 30 nucleotides and most preferably from about 20 to 25 nucleotides. Preferably the fragments can be used in polymerase chain reaction (PCR), various hybridization procedures or microarray procedures to identify or amplify identical or related parts of mRNA or DNA molecules. A fragment or segment may uniquely identify each polynucleotide sequence of the present invention. Preferably the fragment comprises a sequence substantially similar to any one of SEO ID NO: 1-341.

Probes may, for example, be used to determine whether specific mRNA molecules are present in a cell or tissue or to isolate similar nucleic acid sequences from chromosomal DNA as described by Walsh et al. (Walsh, P.S. et al., 1992, PCR Methods Appl 1:241-250). They may be labeled by nick translation, Klenow fill-in reaction, PCR, or other methods well known in the art. Probes of the present invention, their preparation and/or labeling are elaborated in Sambrook, J. et al., 1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, NY; or Ausubel, F.M. et al., 1989, Current Protocols in Molecular

Biology, John Wiley & Sons, New York NY, both of which are incorporated herein by reference in their entirety.

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The nucleic acid sequences of the present invention also include the sequence information from the nucleic acid sequences of SEQ ID NO: 1-341. The sequence information can be a segment of any one of SEQ ID NO: 1-341 that uniquely identifies or represents the sequence information of that sequence of SEQ ID NO: 1-341. One such segment can be a twenty-mer nucleic acid sequence because the probability that a twenty-mer is fully matched in the human genome is 1 in 300. In the human genome, there are three billion base pairs in one set of chromosomes. Because 4²⁰ possible twenty-mers exist, there are 300 times more twenty-mers than there are base pairs in a set of human chromosomes. Using the same analysis, the probability for a seventeen-mer to be fully matched in the human genome is approximately 1 in 5. When these segments are used in arrays for expression studies, fifteen-mer segments can be used. The probability that the fifteen-mer is fully matched in the expressed sequences is also approximately one in five because expressed sequences comprise less than approximately 5% of the entire genome sequence.

Similarly, when using sequence information for detecting a single mismatch, a segment can be a twenty-five mer. The probability that the twenty-five mer would appear in a human genome with a single mismatch is calculated by multiplying the probability for a full match $(1 \div 4^{25})$ times the increased probability for mismatch at each nucleotide position (3 x 25). The probability that an eighteen mer with a single mismatch can be detected in an array for expression studies is approximately one in five. The probability that a twenty-mer with a single mismatch can be detected in a human genome is approximately one in five.

The term "open reading frame," ORF, means a series of nucleotide triplets coding for amino acids without any termination codons and is a sequence translatable into protein.

The terms "operably linked" or "operably associated" refer to functionally related nucleic acid sequences. For example, a promoter is operably associated or operably linked with a coding sequence if the promoter controls the transcription of the coding sequence. While operably linked nucleic acid sequences can be contiguous and in the same reading frame, certain genetic elements e.g. repressor genes are not contiguously linked to the coding sequence but still control transcription/translation of the coding sequence.

The term "pluripotent" refers to the capability of a cell to differentiate into a number of differentiated cell types that are present in an adult organism. A pluripotent cell is restricted in its differentiation capability in comparison to a totipotent cell.

The terms "polypeptide" or "peptide" or "amino acid sequence" refer to an oligopeptide, peptide, polypeptide or protein sequence or fragment thereof and to naturally occurring or synthetic molecules. A polypeptide "fragment," "portion," or "segment" is a stretch of amino acid residues of at least about 5 amino acids, preferably at least about 7 amino acids, more preferably at least about 9 amino acids and most preferably at least about 17 or more amino acids. The peptide preferably is not greater than about 500 amino acids, more preferably less than 200 amino acids more preferably less than 150 amino acids and most preferably less than 100 amino acids. Preferably the peptide is from about 5 to about 200 amino acids. To be active, any polypeptide must have sufficient length to display biological and/or immunological activity.

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The term "naturally occurring polypeptide" refers to polypeptides produced by cells that have not been genetically engineered and specifically contemplates various polypeptides arising from post-translational modifications of the polypeptide including, but not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation and acylation.

The term "translated protein coding portion" means a sequence which encodes for the full length protein which may include any leader sequence or any processing sequence.

The term "mature protein coding sequence" means a sequence which encodes a peptide or protein without a signal or leader sequence. The "mature protein portion" means that portion of the protein which does not include a signal or leader sequence. The peptide may have been produced by processing in the cell which removes any leader/signal sequence. The mature protein portion may or may not include an initial methionine residue. The methionine residue may be removed from the protein during processing in the cell. The peptide may be produced synthetically or the protein may have been produced using a polynucleotide only encoding for the mature protein coding sequence.

The term "derivative" refers to polypeptides chemically modified by such techniques as ubiquitination, labeling (e.g., with radionuclides or various enzymes), covalent polymer attachment such as pegylation (derivatization with polyethylene glycol) and insertion or substitution by chemical synthesis of amino acids such as ornithine, which do not normally occur in human proteins.

The term "variant" (or "analog") refers to any polypeptide differing from naturally occurring polypeptides by amino acid insertions, deletions, and substitutions, created using, e g., recombinant DNA techniques. Guidance in determining which amino acid residues may be replaced, added or deleted without abolishing activities of interest, may be found by

comparing the sequence of the particular polypeptide with that of homologous peptides and minimizing the number of amino acid sequence changes made in regions of high homology (conserved regions) or by replacing amino acids with consensus sequence.

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Alternatively, recombinant variants encoding these same or similar polypeptides may be synthesized or selected by making use of the "redundancy" in the genetic code. Various codon substitutions, such as the silent changes which produce various restriction sites, may be introduced to optimize cloning into a plasmid or viral vector or expression in a particular prokaryotic or eukaryotic system. Mutations in the polynucleotide sequence may be reflected in the polypeptide or domains of other peptides added to the polypeptide to modify the properties of any part of the polypeptide, to change characteristics such as ligand-binding affinities, interchain affinities, or degradation/turnover rate.

Preferably, amino acid "substitutions" are the result of replacing one amino acid with another amino acid having similar structural and/or chemical properties, *i.e.*, conservative amino acid replacements. "Conservative" amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues involved. For example, nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan, and methionine; polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine; positively charged (basic) amino acids include arginine, lysine, and histidine; and negatively charged (acidic) amino acids include aspartic acid and glutamic acid. "Insertions" or "deletions" are preferably in the range of about 1 to 20 amino acids, more preferably 1 to 10 amino acids. The variation allowed may be experimentally determined by systematically making insertions, deletions, or substitutions of amino acids in a polypeptide molecule using recombinant DNA techniques and assaying the resulting recombinant variants for activity.

Alternatively, where alteration of function is desired, insertions, deletions or non-conservative alterations can be engineered to produce altered polypeptides. Such alterations can, for example, alter one or more of the biological functions or biochemical characteristics of the polypeptides of the invention. For example, such alterations may change polypeptide characteristics such as ligand-binding affinities, interchain affinities, or degradation/turnover rate. Further, such alterations can be selected so as to generate polypeptides that are better suited for expression, scale up and the like in the host cells

chosen for expression. For example, cysteine residues can be deleted or substituted with another amino acid residue in order to eliminate disulfide bridges.

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The terms "purified" or "substantially purified" as used herein denotes that the indicated nucleic acid or polypeptide is present in the substantial absence of other biological macromolecules, e.g., polynucleotides, proteins, and the like. In one embodiment, the polynucleotide or polypeptide is purified such that it constitutes at least 95% by weight, more preferably at least 99% by weight, of the indicated biological macromolecules present (but water, buffers, and other small molecules, especially molecules having a molecular weight of less than 1000 daltons, can be present).

The term "isolated" as used herein refers to a nucleic acid or polypeptide separated from at least one other component (e.g., nucleic acid or polypeptide) present with the nucleic acid or polypeptide in its natural source. In one embodiment, the nucleic acid or polypeptide is found in the presence of (if anything) only a solvent, buffer, ion, or other component normally present in a solution of the same. The terms "isolated" and "purified" do not encompass nucleic acids or polypeptides present in their natural source.

The term "recombinant," when used herein to refer to a polypeptide or protein, means that a polypeptide or protein is derived from recombinant (e.g., microbial, insect, or mammalian) expression systems. "Microbial" refers to recombinant polypeptides or proteins made in bacterial or fungal (e.g., yeast) expression systems. As a product, "recombinant microbial" defines a polypeptide or protein essentially free of native endogenous substances and unaccompanied by associated native glycosylation. Polypeptides or proteins expressed in most bacterial cultures, e.g., E. coli, will be free of glycosylation modifications; polypeptides or proteins expressed in yeast will have a glycosylation pattern in general different from those expressed in mammalian cells.

The term "recombinant expression vehicle or vector" refers to a plasmid or phage or virus or vector, for expressing a polypeptide from a DNA (RNA) sequence. An expression vehicle can comprise a transcriptional unit comprising an assembly of (1) a genetic element or elements having a regulatory role in gene expression, for example, promoters or enhancers, (2) a structural or coding sequence which is transcribed into mRNA and translated into protein, and (3) appropriate transcription initiation and termination sequences. Structural units intended for use in yeast or eukaryotic expression systems preferably include a leader sequence enabling extracellular secretion of translated protein by a host cell. Alternatively, where recombinant protein is expressed without a leader or transport sequence, it may include

an amino terminal methionine residue. This residue may or may not be subsequently cleaved from the expressed recombinant protein to provide a final product.

The term "recombinant expression system" means host cells which have stably integrated a recombinant transcriptional unit into chromosomal DNA or carry the recombinant transcriptional unit extrachromosomally. Recombinant expression systems as defined herein will express heterologous polypeptides or proteins upon induction of the regulatory elements linked to the DNA segment or synthetic gene to be expressed. This term also means host cells which have stably integrated a recombinant genetic element or elements having a regulatory role in gene expression, for example, promoters or enhancers. Recombinant expression systems as defined herein will express polypeptides or proteins endogenous to the cell upon induction of the regulatory elements linked to the endogenous DNA segment or gene to be expressed. The cells can be prokaryotic or eukaryotic.

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The term "secreted" includes a protein that is transported across or through a membrane, including transport as a result of signal sequences in its amino acid sequence when it is expressed in a suitable host cell. "Secreted" proteins include without limitation proteins secreted wholly (e.g., soluble proteins) or partially (e.g., receptors) from the cell in which they are expressed. "Secreted" proteins also include without limitation proteins that are transported across the membrane of the endoplasmic reticulum. "Secreted" proteins are also intended to include proteins containing non-typical signal sequences (e.g. Interleukin-1 Beta, see Krasney, P.A. and Young, P.R. (1992) Cytokine 4(2): 134-143) and factors released from damaged cells (e.g. Interleukin-1 Receptor Antagonist, see Arend, W.P. et. al. (1998) Annu. Rev. Immunol. 16:27-55)

Where desired, an expression vector may be designed to contain a "signal or leader sequence" which will direct the polypeptide through the membrane of a cell. Such a sequence may be naturally present on the polypeptides of the present invention or provided from heterologous protein sources by recombinant DNA techniques.

The term "stringent" is used to refer to conditions that are commonly understood in the art as stringent. Stringent conditions can include highly stringent conditions (i.e., hybridization to filter-bound DNA in 0.5 M NaHPO₄, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65°C, and washing in 0.1X SSC/0.1% SDS at 68°C), and moderately stringent conditions (i.e., washing in 0.2X SSC/0.1% SDS at 42°C). Other exemplary hybridization conditions are described herein in the examples.

In instances of hybridization of deoxyoligonucleotides, additional exemplary stringent hybridization conditions include washing in 6X SSC/0.05% sodium pyrophosphate at 37°C (for 14-base oligonucleotides), 48°C (for 17-base oligos), 55°C (for 20-base oligonucleotides), and 60°C (for 23-base oligonucleotides).

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As used herein, "substantially equivalent" can refer both to nucleotide and amino acid sequences, for example a mutant sequence, that varies from a reference sequence by one or more substitutions, deletions, or additions, the net effect of which does not result in an adverse functional dissimilarity between the reference and subject sequences. Typically, such a substantially equivalent sequence varies from one of those listed herein by no more than about 35% (i.e., the number of individual residue substitutions, additions, and/or deletions in a substantially equivalent sequence, as compared to the corresponding reference sequence, divided by the total number of residues in the substantially equivalent sequence is about 0.35 or less). Such a sequence is said to have 65% sequence identity to the listed sequence. In one embodiment, a substantially equivalent, e.g., mutant, sequence of the invention varies from a listed sequence by no more than 30% (70% sequence identity); in a variation of this embodiment, by no more than 25% (75% sequence identity); and in a further variation of this embodiment, by no more than 20% (80% sequence identity) and in a further variation of this embodiment, by no more than 10% (90% sequence identity) and in a further variation of this embodiment, by no more that 5% (95% sequence identity). Substantially equivalent, e.g., mutant, amino acid sequences according to the invention preferably have at least 80% sequence identity with a listed amino acid sequence, more preferably at least 85% sequence identity, more preferably at least 90% sequence identity, more preferably at least 95% identity, more preferably at least 98% identity, and most preferably at least 99% identity. Substantially equivalent nucleotide sequences of the invention can have lower percent sequence identities, taking into account, for example, the redundancy or degeneracy of the genetic code. Preferably, nucleotide sequence has at least about 65% identity, more preferably at least about 75% identity, more preferably at least about 80% sequence identity, more preferably at least about 85% sequence identity, more preferably at least about 90% sequence identity, and most preferably at least about 95% identity, more preferably at least about 98% sequence identity, and most preferably at least about 99% sequence identity. For the purposes of the present invention, sequences having substantially equivalent biological activity and substantially equivalent expression characteristics are considered substantially equivalent. For the purposes of determining equivalence, truncation of the mature sequence

(e.g., via a mutation which creates a spurious stop codon) should be disregarded. Sequence identity may be determined, e.g., using the Jotun Hein method (Hein, J. (1990) Methods Enzymol. 183:626-645). Identity between sequences can also be determined by other methods known in the art, e.g. by varying hybridization conditions.

The term "totipotent" refers to the capability of a cell to differentiate into all of the cell types of an adult organism.

The term "transformation" means introducing DNA into a suitable host cell so that the DNA is replicable, either as an extrachromosomal element, or by chromosomal integration. The term "transfection" refers to the taking up of an expression vector by a suitable host cell, whether or not any coding sequences are in fact expressed. The term "infection" refers to the introduction of nucleic acids into a suitable host cell by use of a virus or viral vector.

As used herein, an "uptake modulating fragment," UMF, means a series of nucleotides which mediate the uptake of a linked DNA fragment into a cell. UMFs can be readily identified using known UMFs as a target sequence or target motif with the computer-based systems described below. The presence and activity of a UMF can be confirmed by attaching the suspected UMF to a marker sequence. The resulting nucleic acid molecule is then incubated with an appropriate host under appropriate conditions and the uptake of the marker sequence is determined. As described above, a UMF will increase the frequency of uptake of a linked marker sequence.

Each of the above terms is meant to encompass all that is described for each, unless the context dictates otherwise.

4.2 NUCLEIC ACIDS OF THE INVENTION

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Nucleotide sequences of the invention are set forth in the Sequence Listing.

The isolated polynucleotides of the invention include a polynucleotide comprising the nucleotide sequences of SEQ ID NO: 1-341; a polynucleotide encoding any one of the peptide sequences of SEQ ID NO: 342-682; and a polynucleotide comprising the nucleotide sequence encoding the mature protein coding sequence of the polypeptides of any one of SEQ ID NO: 342-682. The polynucleotides of the present invention also include, but are not limited to, a polynucleotide that hybridizes under stringent conditions to (a) the complement of any of the nucleotides sequences of SEQ ID NO: 1-341; (b) nucleotide sequences encoding any one of the amino acid sequences set forth in the Sequence Listing as SEQ ID NO: 342-682; (c) a polynucleotide which is an allelic variant of any polynucleotide recited above; (d)

a polynucleotide which encodes a species homolog of any of the proteins recited above; or (e) a polynucleotide that encodes a polypeptide comprising a specific domain or truncation of the polypeptides of SEQ ID NO: 342-682. Domains of interest may depend on the nature of the encoded polypeptide; e.g., domains in receptor-like polypeptides include ligand-binding, extracellular, transmembrane, or cytoplasmic domains, or combinations thereof; domains in immunoglobulin-like proteins include the variable immunoglobulin-like domains; domains in enzyme-like polypeptides include catalytic and substrate binding domains; and domains in ligand polypeptides include receptor-binding domains.

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The polynucleotides of the invention include naturally occurring or wholly or partially synthetic DNA, e.g., cDNA and genomic DNA, and RNA, e.g., mRNA. The polynucleotides may include all of the coding region of the cDNA or may represent a portion of the coding region of the cDNA.

The present invention also provides genes corresponding to the cDNA sequences disclosed herein. The corresponding genes can be isolated in accordance with known methods using the sequence information disclosed herein. Such methods include the preparation of probes or primers from the disclosed sequence information for identification and/or amplification of genes in appropriate genomic libraries or other sources of genomic materials. Further 5' and 3' sequence can be obtained using methods known in the art. For example, full length cDNA or genomic DNA that corresponds to any of the polynucleotides of SEQ ID NO: 1-341 can be obtained by screening appropriate cDNA or genomic DNA libraries under suitable hybridization conditions using any of the polynucleotides of SEQ ID NO: 1-341 or a portion thereof as a probe. Alternatively, the polynucleotides of SEQ ID NO: 1-341 may be used as the basis for suitable primer(s) that allow identification and/or amplification of genes in appropriate genomic DNA or cDNA libraries.

The nucleic acid sequences of the invention can be assembled from ESTs and sequences (including cDNA and genomic sequences) obtained from one or more public databases, such as dbEST, gbpri, and UniGene. The EST sequences can provide identifying sequence information, representative fragment or segment information, or novel segment information for the full-length gene.

The polynucleotides of the invention also provide polynucleotides including nucleotide sequences that are substantially equivalent to the polynucleotides recited above. Polynucleotides according to the invention can have, e.g., at least about 65%, at least about 70%, at least about 75%, at least about 80%, 81%, 82%, 83%, 84%, more typically at least

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about 85%, 86%, 87%, 88%, 89%, more typically at least about 90%, 91%, 92%, 93%, 94%, and even more typically at least about 95%, 96%, 97%, 98%, 99%, sequence identity to a polynucleotide recited above.

Included within the scope of the nucleic acid sequences of the invention are nucleic acid sequence fragments that hybridize under stringent conditions to any of the nucleotide sequences of SEQ ID NO: 1-341, or complements thereof, which fragment is greater than about 5 nucleotides, preferably 7 nucleotides, more preferably greater than 9 nucleotides and most preferably greater than 17 nucleotides. Fragments of, e.g. 15, 17, or 20 nucleotides or more that are selective for (i.e. specifically hybridize to) any one of the polynucleotides of the invention are contemplated. Probes capable of specifically hybridizing to a polynucleotide can differentiate polynucleotide sequences of the invention from other polynucleotide sequences in the same family of genes or can differentiate human genes from genes of other species, and are preferably based on unique nucleotide sequences.

The sequences falling within the scope of the present invention are not limited to these specific sequences, but also include allelic and species variations thereof. Allelic and species variations can be routinely determined by comparing the sequence provided in SEQ ID NO: 1-341, a representative fragment thereof, or a nucleotide sequence at least 90% identical, preferably 95% identical, to SEQ ID NO: 1-341 with a sequence from another isolate of the same species. Furthermore, to accommodate codon variability, the invention includes nucleic acid molecules coding for the same amino acid sequences as do the specific ORFs disclosed herein. In other words, in the coding region of an ORF, substitution of one codon for another codon that encodes the same amino acid is expressly contemplated.

The nearest neighbor or homology result for the nucleic acids of the present invention, including SEQ ID NO: 1-341, can be obtained by searching a database using an algorithm or a program. Preferably, a BLAST which stands for Basic Local Alignment Search Tool is used to search for local sequence alignments (Altshul, S.F. J Mol. Evol. 36 290-300 (1993) and Altschul S.F. et al. J. Mol. Biol. 21:403-410 (1990)). Alternatively a FASTA version 3 search against Genpept, using Fastxy algorithm.

Species homologs (or orthologs) of the disclosed polynucleotides and proteins are also provided by the present invention. Species homologs may be isolated and identified by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source from the desired species.

The invention also encompasses allelic variants of the disclosed polynucleotides or proteins; that is, naturally-occurring alternative forms of the isolated polynucleotide which also encode proteins which are identical, homologous or related to that encoded by the polynucleotides.

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The nucleic acid sequences of the invention are further directed to sequences which encode variants of the described nucleic acids. These amino acid sequence variants may be prepared by methods known in the art by introducing appropriate nucleotide changes into a native or variant polynucleotide. There are two variables in the construction of amino acid sequence variants: the location of the mutation and the nature of the mutation. Nucleic acids encoding the amino acid sequence variants are preferably constructed by mutating the polynucleotide to encode an amino acid sequence that does not occur in nature. These nucleic acid alterations can be made at sites that differ in the nucleic acids from different species (variable positions) or in highly conserved regions (constant regions). Sites at such locations will typically be modified in series, e.g., by substituting first with conservative choices (e.g., hydrophobic amino acid to a different hydrophobic amino acid) and then with more distant choices (e.g., hydrophobic amino acid to a charged amino acid), and then deletions or insertions may be made at the target site. Amino acid sequence deletions generally range from about 1 to 30 residues, preferably about 1 to 10 residues, and are typically contiguous. Amino acid insertions include amino- and/or carboxyl-terminal fusions ranging in length from one to one hundred or more residues, as well as intrasequence insertions of single or multiple amino acid residues. Intrasequence insertions may range generally from about 1 to 10 amino residues, preferably from 1 to 5 residues. Examples of terminal insertions include the heterologous signal sequences necessary for secretion or for intracellular targeting in different host cells and sequences such as FLAG or poly-histidine sequences useful for purifying the expressed protein.

In a preferred method, polynucleotides encoding the novel amino acid sequences are changed via site-directed mutagenesis. This method uses oligonucleotide sequences to alter a polynucleotide to encode the desired amino acid variant, as well as sufficient adjacent nucleotides on both sides of the changed amino acid to form a stable duplex on either side of the site of being changed. In general, the techniques of site-directed mutagenesis are well known to those of skill in the art and this technique is exemplified by publications such as, Edelman et al., *DNA* 2:183 (1983). A versatile and efficient method for producing site-specific changes in a polynucleotide sequence was published by Zoller and Smith,

Nucleic Acids Res. 10:6487-6500 (1982). PCR may also be used to create amino acid sequence variants of the novel nucleic acids. When small amounts of template DNA are used as starting material, primer(s) that differs slightly in sequence from the corresponding region in the template DNA can generate the desired amino acid variant. PCR amplification results in a population of product DNA fragments that differ from the polynucleotide template encoding the polypeptide at the position specified by the primer. The product DNA fragments replace the corresponding region in the plasmid and this gives a polynucleotide encoding the desired amino acid variant.

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A further technique for generating amino acid variants is the cassette mutagenesis technique described in Wells et al., *Gene* 34:315 (1985); and other mutagenesis techniques well known in the art, such as, for example, the techniques in Sambrook et al., supra, and *Current Protocols in Molecular Biology*, Ausubel et al. Due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the same or a functionally equivalent amino acid sequence may be used in the practice of the invention for the cloning and expression of these novel nucleic acids. Such DNA sequences include those which are capable of hybridizing to the appropriate novel nucleic acid sequence under stringent conditions.

Polynucleotides encoding preferred polypeptide truncations of the invention can be used to generate polynucleotides encoding chimeric or fusion proteins comprising one or more domains of the invention and heterologous protein sequences.

The polynucleotides of the invention additionally include the complement of any of the polynucleotides recited above. The polynucleotide can be DNA (genomic, cDNA, amplified, or synthetic) or RNA. Methods and algorithms for obtaining such polynucleotides are well known to those of skill in the art and can include, for example, methods for determining hybridization conditions that can routinely isolate polynucleotides of the desired sequence identities.

In accordance with the invention, polynucleotide sequences comprising the mature protein coding sequences corresponding to any one of SEQ ID NO: 1-341, or functional equivalents thereof, may be used to generate recombinant DNA molecules that direct the expression of that nucleic acid, or a functional equivalent thereof, in appropriate host cells. Also included are the cDNA inserts of any of the clones identified herein.

A polynucleotide according to the invention can be joined to any of a variety of other nucleotide sequences by well-established recombinant DNA techniques (see Sambrook J et

al. (1989) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, NY). Useful nucleotide sequences for joining to polynucleotides include an assortment of vectors, e.g., plasmids, cosmids, lambda phage derivatives, phagemids, and the like, that are well known in the art. Accordingly, the invention also provides a vector including a polynucleotide of the invention and a host cell containing the polynucleotide. In general, the vector contains an origin of replication functional in at least one organism, convenient restriction endonuclease sites, and a selectable marker for the host cell. Vectors according to the invention include expression vectors, replication vectors, probe generation vectors, and sequencing vectors. A host cell according to the invention can be a prokaryotic or eukaryotic cell and can be a unicellular organism or part of a multicellular organism.

The present invention further provides recombinant constructs comprising a nucleic acid having any of the nucleotide sequences of SEQ ID NO: 1-341 or a fragment thereof or any other polynucleotides of the invention. In one embodiment, the recombinant constructs of the present invention comprise a vector, such as a plasmid or viral vector, into which a nucleic acid having any of the nucleotide sequences of SEQ ID NO: 1-341 or a fragment thereof is inserted, in a forward or reverse orientation. In the case of a vector comprising one of the ORFs of the present invention, the vector may further comprise regulatory sequences, including for example, a promoter, operably linked to the ORF. Large numbers of suitable vectors and promoters are known to those of skill in the art and are commercially available for generating the recombinant constructs of the present invention. The following vectors are provided by way of example. Bacterial: pBs, phagescript, PsiX174, pBluescript SK, pBs KS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene); pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia). Eukaryotic: pWLneo, pSV2cat, pOG44, PXTI, pSG (Stratagene) pSVK3, pBPV, pMSG, pSVL (Pharmacia).

The isolated polynucleotide of the invention may be operably linked to an expression control sequence such as the pMT2 or pED expression vectors disclosed in Kaufman et al., Nucleic Acids Res. 19, 4485-4490 (1991), in order to produce the protein recombinantly. Many suitable expression control sequences are known in the art. General methods of expressing recombinant proteins are also known and are exemplified in R. Kaufman, Methods in Enzymology 185, 537-566 (1990). As defined herein "operably linked" means that the isolated polynucleotide of the invention and an expression control sequence are situated within a vector or cell in such a way that the protein is expressed by a host cell which has been transformed (transfected) with the ligated polynucleotide/expression control sequence.

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Promoter regions can be selected from any desired gene using CAT (chloramphenicol transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda PR, and trc. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art. Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, e.g., the ampicillin resistance gene of E. coli and S. cerevisiae TRP1 gene, and a promoter derived from a highly-expressed gene to direct transcription of a downstream structural sequence. Such promoters can be derived from operons encoding glycolytic enzymes such as 3-phosphoglycerate kinase (PGK), a-factor, acid phosphatase, or heat shock proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences, and preferably, a leader sequence capable of directing secretion of translated protein into the periplasmic space or extracellular medium. Optionally, the heterologous sequence can encode a fusion protein including an amino terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product. Useful expression vectors for bacterial use are constructed by inserting a structural DNA sequence encoding a desired protein together with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector and to, if desirable, provide amplification within the host. Suitable prokaryotic hosts for transformation include E. coli, Bacillus subtilis, Salmonella typhimurium and various species within the genera Pseudomonas, Streptomyces, and Staphylococcus, although others may also be employed as a matter of choice.

As a representative but non-limiting example, useful expression vectors for bacterial use can comprise a selectable marker and bacterial origin of replication derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and GEM 1 (Promega Biotech, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed. Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter is induced

or derepressed by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period. Cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification.

Polynucleotides of the invention can also be used to induce immune responses. For example, as described in Fan et al., *Nat. Biotech.* 17:870-872 (1999), incorporated herein by reference, nucleic acid sequences encoding a polypeptide may be used to generate antibodies against the encoded polypeptide following topical administration of naked plasmid DNA or following injection, and preferably intramuscular injection of the DNA. The nucleic acid sequences are preferably inserted in a recombinant expression vector and may be in the form of naked DNA.

4.3 ANTISENSE NUCLEIC ACIDS

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Another aspect of the invention pertains to isolated antisense nucleic acid molecules that are hybridizable to or complementary to the nucleic acid molecule comprising the nucleotide sequence of SEQ ID NO: 1-341, or fragments, analogs or derivatives thereof. An "antisense" nucleic acid comprises a nucleotide sequence that is complementary to a "sense" nucleic acid encoding a protein, e.g., complementary to the coding strand of a double-stranded cDNA molecule or complementary to an mRNA sequence. In specific aspects, antisense nucleic acid molecules are provided that comprise a sequence complementary to at least about 10, 25, 50, 100, 250 or 500 nucleotides or an entire coding strand, or to only a portion thereof. Nucleic acid molecules encoding fragments, homologs, derivatives and analogs of a protein of any of SEQ ID NO: 342-682 or antisense nucleic acids complementary to a nucleic acid sequence of SEQ ID NO: 1-341 are additionally provided.

In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence of the invention. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid residues. In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence of the invention. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (i.e., also referred to as 5' and 3' untranslated regions).

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Given the coding strand sequences encoding a nucleic acid disclosed herein (e.g., SEQ ID NO: 1-341), antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick or Hoogsteen base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of an mRNA, but more preferably is an oligonucleotide that is antisense to only a portion of the coding or noncoding region of a mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of a mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length. An antisense nucleic acid of the invention can be constructed using chemical synthesis or enzymatic ligation reactions using procedures known in the art. For example, an antisense nucleic acid (e.g., an antisense oligonucleotide) can be chemically synthesized using naturally occurring nucleotides or variously modified nucleotides designed to increase the biological stability of the molecules or to increase the physical stability of the duplex formed between the antisense and sense nucleic acids, e.g., phosphorothioate derivatives and acridine substituted nucleotides can be used.

Examples of modified nucleotides that can be used to generate the antisense nucleic acid include: 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine. pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v). 5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine. Alternatively, the antisense nucleic acid can be produced biologically using an expression vector into which a nucleic acid has been subcloned in an antisense orientation (i.e., RNA transcribed from the inserted nucleic acid will be of an antisense orientation to a target nucleic acid of interest, described further in the following subsection).

The antisense nucleic acid molecules of the invention are typically administered to a subject or generated *in situ* such that they hybridize with or bind to cellular mRNA and/or

genomic DNA encoding a protein according to the invention to thereby inhibit expression of the protein, e.g., by inhibiting transcription and/or translation. The hybridization can be by conventional nucleotide complementarity to form a stable duplex, or, for example, in the case of an antisense nucleic acid molecule that binds to DNA duplexes, through specific interactions in the major groove of the double helix. An example of a route of administration of antisense nucleic acid molecules of the invention includes direct injection at a tissue site. Alternatively, antisense nucleic acid molecules can be modified to target selected cells and then administered systemically. For example, for systemic administration, antisense molecules can be modified such that they specifically bind to receptors or antigens expressed on a selected cell surface, e.g., by linking the antisense nucleic acid molecules to peptides or antibodies that bind to cell surface receptors or antigens. The antisense nucleic acid molecules can also be delivered to cells using the vectors described herein. To achieve sufficient intracellular concentrations of antisense molecules, vector constructs in which the antisense nucleic acid molecule is placed under the control of a strong pol II or pol III promoter are preferred.

In yet another embodiment, the antisense nucleic acid molecule of the invention is an α-anomeric nucleic acid molecule. An α-anomeric nucleic acid molecule forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual β-units, the strands run parallel to each other (Gaultier et al. (1987) Nucleic Acids Res 15: 6625-6641). The antisense nucleic acid molecule can also comprise a 2'-o-methylribonucleotide (Inoue et al. (1987) Nucleic Acids Res 15: 6131-6148) or a chimeric RNA -DNA analogue (Inoue et al. (1987) FEBS Lett 215: 327-330).

4.4 RIBOZYMES AND PNA MOIETIES

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In still another embodiment, an antisense nucleic acid of the invention is a ribozyme. Ribozymes are catalytic RNA molecules with ribonuclease activity that are capable of cleaving a single-stranded nucleic acid, such as an mRNA, to which they have a complementary region. Thus, ribozymes (e.g., hammerhead ribozymes (described in Haselhoff and Gerlach (1988) Nature 334:585-591)) can be used to catalytically cleave a mRNA transcripts to thereby inhibit translation of a mRNA. A ribozyme having specificity for a nucleic acid of the invention can be designed based upon the nucleotide sequence of a DNA disclosed herein (i.e., SEQ ID NO: 1-341). For example, a derivative of a Tetrahymena L-19 IVS RNA can be constructed in which the nucleotide sequence of the active site is

complementary to the nucleotide sequence to be cleaved in an mRNA of SEQ ID NO: 1-341 (see, e.g., Cech et al. U.S. Pat. No. 4,987,071; and Cech et al. U.S. Pat. No. 5,116,742). Alternatively, polynucleotides of the invention can be used to select a catalytic RNA having a specific ribonuclease activity from a pool of RNA molecules. See, e.g., Bartel et al., (1993) Science 261:1411-1418.

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Alternatively, gene expression can be inhibited by targeting nucleotide sequences complementary to the regulatory region (e.g., promoter and/or enhancers) to form triple helical structures that prevent transcription of the gene in target cells. See generally, Helene. (1991) Anticancer Drug Des. 6: 569-84; Helene. et al. (1992) Ann. N.Y. Acad. Sci. 660:27-36; and Maher (1992) Bioassays 14: 807-15.

In various embodiments, the nucleic acids of the invention can be modified at the base moiety, sugar moiety or phosphate backbone to improve, e.g., the stability, hybridization, or solubility of the molecule. For example, the deoxyribose phosphate backbone of the nucleic acids can be modified to generate peptide nucleic acids (see Hyrup et al. (1996) Bioorg Med Chem 4: 5-23). As used herein, the terms "peptide nucleic acids" or "PNAs" refer to nucleic acid mimics, e.g., DNA mimics, in which the deoxyribose phosphate backbone is replaced by a pseudopeptide backbone and only the four natural nucleobases are retained. The neutral backbone of PNAs has been shown to allow for specific hybridization to DNA and RNA under conditions of low ionic strength. The synthesis of PNA oligomers can be performed using standard solid phase peptide synthesis protocols as described in Hyrup et al. (1996) above; Perry-O'Keefe et al. (1996) PNAS 93: 14670-675.

PNAs of the invention can be used in therapeutic and diagnostic applications. For example, PNAs can be used as antisense or antigene agents for sequence-specific modulation of gene expression by, e.g., inducing transcription or translation arrest or inhibiting replication. PNAs of the invention can also be used, e.g., in the analysis of single base pair mutations in a gene by, e.g., PNA directed PCR clamping; as artificial restriction enzymes when used in combination with other enzymes, e.g., S1 nucleases (Hyrup B. (1996) above); or as probes or primers for DNA sequence and hybridization (Hyrup et al. (1996), above; Perry-O'Keefe (1996), above).

In another embodiment, PNAs of the invention can be modified, e.g., to enhance their stability or cellular uptake, by attaching lipophilic or other helper groups to PNA, by the formation of PNA-DNA chimeras, or by the use of liposomes or other techniques of drug delivery known in the art. For example, PNA-DNA chimeras can be generated that may

combine the advantageous properties of PNA and DNA. Such chimeras allow DNA recognition enzymes, e.g., RNase H and DNA polymerases, to interact with the DNA portion while the PNA portion would provide high binding affinity and specificity. PNA-DNA chimeras can be linked using linkers of appropriate lengths selected in terms of base stacking, number of bonds between the nucleobases, and orientation (Hyrup (1996) above). The synthesis of PNA-DNA chimeras can be performed as described in Hyrup (1996) above and Finn et al. (1996) Nucl Acids Res 24: 3357-63. For example, a DNA chain can be synthesized on a solid support using standard phosphoramidite coupling chemistry, and modified nucleoside analogs, e.g., 5'-(4-methoxytrityl)amino-5'-deoxy-thymidine phosphoramidite, can be used between the PNA and the 5' end of DNA (Mag et al. (1989) Nucl Acid Res 17: 5973-88). PNA monomers are then coupled in a stepwise manner to produce a chimeric molecule with a 5' PNA segment and a 3' DNA segment (Finn et al. (1996) above). Alternatively, chimeric molecules can be synthesized with a 5' DNA segment and a 3' PNA segment. See, Petersen et al. (1975) Bioorg Med Chem Lett 5: 1119-11124.

In other embodiments, the oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. W088/09810) or the blood-brain barrier (see, e.g., PCT Publication No. W089/10134). In addition, oligonucleotides can be modified with hybridization triggered cleavage agents (See, e.g., Krol et al., 1988, BioTechniques 6:958-976) or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5: 539-549). To this end, the oligonucleotide may be conjugated to another molecule, e.g., a peptide, a hybridization triggered cross-linking agent, a transport agent, a hybridization-triggered cleavage agent, etc.

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4.5 HOSTS

The present invention further provides host cells genetically engineered to contain the polynucleotides of the invention. For example, such host cells may contain nucleic acids of the invention introduced into the host cell using known transformation, transfection or infection methods. The present invention still further provides host cells genetically engineered to express the polynucleotides of the invention, wherein such polynucleotides are in operative association with a regulatory sequence heterologous to the host cell which drives expression of the polynucleotides in the cell.

Knowledge of nucleic acid sequences allows for modification of cells to permit, or increase, expression of endogenous polypeptide. Cells can be modified (e.g., by homologous recombination) to provide increased polypeptide expression by replacing, in whole or in part, the naturally occurring promoter with all or part of a heterologous promoter so that the cells express the polypeptide at higher levels. The heterologous promoter is inserted in such a manner that it is operatively linked to the encoding sequences. See, for example, PCT International Publication No. WO94/12650, PCT International Publication No. WO92/20808, and PCT International Publication No. WO91/09955. It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (e.g., ada, dhfr, and the multifunctional CAD gene which encodes carbamyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the coding sequence, amplification of the marker DNA by standard selection methods results in co-amplification of the desired protein coding sequences in the cells.

The host cell can be a higher eukaryotic host cell, such as a mammalian cell, a lower eukaryotic host cell, such as a yeast cell, or the host cell can be a prokaryotic cell, such as a bacterial cell. Introduction of the recombinant construct into the host cell can be effected by calcium phosphate transfection, DEAE-dextran mediated transfection, or electroporation (Davis, L. et al., *Basic Methods in Molecular Biology* (1986)). The host cells containing one of the polynucleotides of the invention, can be used in conventional manners to produce the gene product encoded by the isolated fragment (in the case of an ORF) or can be used to produce a heterologous protein under the control of the EMF.

Any host/vector system can be used to express one or more of the ORFs of the present invention. These include, but are not limited to, eukaryotic hosts such as HeLa cells, Cv-1 cell, COS cells, 293 cells, and Sf9 cells, as well as prokaryotic host such as E. coli and B. subtilis. The most preferred cells are those which do not normally express the particular polypeptide or protein or which expresses the polypeptide or protein at low natural level. Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described by Sambrook, et al., in Molecular Cloning: A Laboratory Manual, Second Edition,

Cold Spring Harbor, New York (1989), the disclosure of which is hereby incorporated by reference.

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Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described by Gluzman, Cell 23:175 (1981). Other cell lines capable of expressing a compatible vector are, for example, the C127, monkey COS cells, Chinese Hamster Ovary (CHO) cells, human kidney 293 cells, human epidermal A431 cells, human Colo205 cells, 3T3 cells, CV-1 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from in vitro culture of primary tissue, primary explants, HeLa cells, mouse L cells, BHK, HL-60, U937, HaK or Jurkat cells. Mammalian expression vectors will comprise an origin of replication, a suitable promoter and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 viral genome, for example, SV40 origin, early promoter, enhancer, splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements. Recombinant polypeptides and proteins produced in bacterial culture are usually isolated by initial extraction from cell pellets, followed by one or more salting-out, aqueous ion exchange or size exclusion chromatography steps. Protein refolding steps can be used, as necessary, in completing configuration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps. Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents.

Alternatively, it may be possible to produce the protein in lower eukaryotes such as yeast or insects or in prokaryotes such as bacteria. Potentially suitable yeast strains include Saccharomyces cerevisiae, Schizosaccharomyces pombe, Kluyveromyces strains, Candida, or any yeast strain capable of expressing heterologous proteins. Potentially suitable bacterial strains include Escherichia coli, Bacillus subtilis, Salmonella typhimurium, or any bacterial strain capable of expressing heterologous proteins. If the protein is made in yeast or bacteria, it may be necessary to modify the protein produced therein, for example by phosphorylation or glycosylation of the appropriate sites, in order to obtain the functional protein. Such covalent attachments may be accomplished using known chemical or enzymatic methods.

In another embodiment of the present invention, cells and tissues may be engineered to express an endogenous gene comprising the polynucleotides of the invention under the

control of inducible regulatory elements, in which case the regulatory sequences of the endogenous gene may be replaced by homologous recombination. As described herein, gene targeting can be used to replace a gene's existing regulatory region with a regulatory sequence isolated from a different gene or a novel regulatory sequence synthesized by genetic engineering methods. Such regulatory sequences may be comprised of promoters, enhancers, scaffold-attachment regions, negative regulatory elements, transcriptional initiation sites, regulatory protein binding sites or combinations of said sequences. Alternatively, sequences which affect the structure or stability of the RNA or protein produced may be replaced, removed, added, or otherwise modified by targeting. These sequence include polyadenylation signals, mRNA stability elements, splice sites, leader sequences for enhancing or modifying transport or secretion properties of the protein, or other sequences which alter or improve the function or stability of protein or RNA molecules.

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The targeting event may be a simple insertion of the regulatory sequence, placing the gene under the control of the new regulatory sequence, e.g., inserting a new promoter or enhancer or both upstream of a gene. Alternatively, the targeting event may be a simple deletion of a regulatory element, such as the deletion of a tissue-specific negative regulatory element. Alternatively, the targeting event may replace an existing element; for example, a tissue-specific enhancer can be replaced by an enhancer that has broader or different cell-type specificity than the naturally occurring elements. Here, the naturally occurring sequences are deleted and new sequences are added. In all cases, the identification of the targeting event may be facilitated by the use of one or more selectable marker genes that are contiguous with the targeting DNA, allowing for the selection of cells in which the exogenous DNA has integrated into the host cell genome. The identification of the targeting event may also be facilitated by the use of one or more marker genes exhibiting the property of negative selection, such that the negatively selectable marker is linked to the exogenous DNA, but configured such that the negatively selectable marker flanks the targeting sequence, and such that a correct homologous recombination event with sequences in the host cell genome does not result in the stable integration of the negatively selectable marker. Markers useful for this purpose include the Herpes Simplex Virus thymidine kinase (TK) gene or the bacterial xanthine-guanine phosphoribosyl-transferase (gpt) gene.

The gene targeting or gene activation techniques which can be used in accordance with this aspect of the invention are more particularly described in U.S. Patent No. 5,272,071 to Chappel; U.S. Patent No. 5,578,461 to Sherwin et al.; International Application No.

PCT/US92/09627 (WO93/09222) by Selden et al.; and International Application No. PCT/US90/06436 (WO91/06667) by Skoultchi et al., each of which is incorporated by reference herein in its entirety.

4.6 POLYPEPTIDES OF THE INVENTION

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The isolated polypeptides of the invention include, but are not limited to, a polypeptide comprising: the amino acid sequences set forth as any one of SEQ ID NO: 342-682 or an amino acid sequence encoded by any one of the nucleotide sequences SEQ ID NO: 1-341 or the corresponding full length or mature protein. Polypeptides of the invention also include polypeptides preferably with biological or immunological activity that are encoded by: (a) a polynucleotide having any one of the nucleotide sequences set forth in SEQ ID NO: 1-341 or (b) polynucleotides encoding any one of the amino acid sequences set forth as SEQ ID NO: 342-682 or (c) polynucleotides that hybridize to the complement of the polynucleotides of either (a) or (b) under stringent hybridization conditions. The invention also provides biologically active or immunologically active variants of any of the amino acid sequences set forth as SEQ ID NO: 342-682 or the corresponding full length or mature protein; and "substantial equivalents" thereof (e.g., with at least about 65%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, 86%, 87%, 88%, 89%, at least about 90%, 91%, 92%, 93%, 94%, typically at least about 95%, 96%, 97%, more typically at least about 98%, or most typically at least about 99% amino acid identity) that retain biological activity. Polypeptides encoded by allelic variants may have a similar. increased, or decreased activity compared to polypeptides comprising SEQ ID NO: 342-682.

Fragments of the proteins of the present invention which are capable of exhibiting biological activity are also encompassed by the present invention. Fragments of the protein may be in linear form or they may be cyclized using known methods, for example, as described in H. U. Saragovi, et al., Bio/Technology 10, 773-778 (1992) and in R. S. McDowell, et al., J. Amer. Chem. Soc. 114, 9245-9253 (1992), both of which are incorporated herein by reference. Such fragments may be fused to carrier molecules such as immunoglobulins for many purposes, including increasing the valency of protein binding sites.

The present invention also provides both full-length and mature forms (for example, without a signal sequence or precursor sequence) of the disclosed proteins. The protein coding sequence is identified in the sequence listing by translation of the disclosed nucleotide

sequences. The mature form of such protein may be obtained by expression of a full-length polynucleotide in a suitable mammalian cell or other host cell. The sequence of the mature form of the protein is also determinable from the amino acid sequence of the full-length form. Where proteins of the present invention are membrane bound, soluble forms of the proteins are also provided. In such forms, part or all of the regions causing the proteins to be membrane bound are deleted so that the proteins are fully secreted from the cell in which they are expressed.

Protein compositions of the present invention may further comprise an acceptable carrier, such as a hydrophilic, e.g., pharmaceutically acceptable, carrier.

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The present invention further provides isolated polypeptides encoded by the nucleic acid fragments of the present invention or by degenerate variants of the nucleic acid fragments of the present invention. By "degenerate variant" is intended nucleotide fragments which differ from a nucleic acid fragment of the present invention (e.g., an ORF) by nucleotide sequence but, due to the degeneracy of the genetic code, encode an identical polypeptide sequence. Preferred nucleic acid fragments of the present invention are the ORFs that encode proteins.

A variety of methodologies known in the art can be utilized to obtain any one of the isolated polypeptides or proteins of the present invention. At the simplest level, the amino acid sequence can be synthesized using commercially available peptide synthesizers. The synthetically-constructed protein sequences, by virtue of sharing primary, secondary or tertiary structural and/or conformational characteristics with proteins may possess biological properties in common therewith, including protein activity. This technique is particularly useful in producing small peptides and fragments of larger polypeptides. Fragments are useful, for example, in generating antibodies against the native polypeptide. Thus, they may be employed as biologically active or immunological substitutes for natural, purified proteins in screening of therapeutic compounds and in immunological processes for the development of antibodies.

The polypeptides and proteins of the present invention can alternatively be purified from cells which have been altered to express the desired polypeptide or protein. As used herein, a cell is said to be altered to express a desired polypeptide or protein when the cell, through genetic manipulation, is made to produce a polypeptide or protein which it normally does not produce or which the cell normally produces at a lower level. One skilled in the art can readily adapt procedures for introducing and expressing either recombinant or synthetic

sequences into eukaryotic or prokaryotic cells in order to generate a cell which produces one of the polypeptides or proteins of the present invention.

The invention also relates to methods for producing a polypeptide comprising growing a culture of host cells of the invention in a suitable culture medium, and purifying the protein from the cells or the culture in which the cells are grown. For example, the methods of the invention include a process for producing a polypeptide in which a host cell containing a suitable expression vector that includes a polynucleotide of the invention is cultured under conditions that allow expression of the encoded polypeptide. The polypeptide can be recovered from the culture, conveniently from the culture medium, or from a lysate prepared from the host cells and further purified. Preferred embodiments include those in which the protein produced by such process is a full length or mature form of the protein.

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In an alternative method, the polypeptide or protein is purified from bacterial cells which naturally produce the polypeptide or protein. One skilled in the art can readily follow known methods for isolating polypeptides and proteins in order to obtain one of the isolated polypeptides or proteins of the present invention. These include, but are not limited to, immunochromatography, HPLC, size-exclusion chromatography, ion-exchange chromatography, and immuno-affinity chromatography. See, e.g., Scopes, Protein Purification: Principles and Practice, Springer-Verlag (1994); Sambrook, et al., in Molecular Cloning: A Laboratory Manual; Ausubel et al., Current Protocols in Molecular Biology. Polypeptide fragments that retain biological/immunological activity include fragments comprising greater than about 100 amino acids, or greater than about 200 amino acids, and fragments that encode specific protein domains.

The purified polypeptides can be used in *in vitro* binding assays which are well known in the art to identify molecules which bind to the polypeptides. These molecules include but are not limited to, for e.g., small molecules, molecules from combinatorial libraries, antibodies or other proteins. The molecules identified in the binding assay are then tested for antagonist or agonist activity in *in vivo* tissue culture or animal models that are well known in the art. In brief, the molecules are titrated into a plurality of cell cultures or animals and then tested for either cell/animal death or prolonged survival of the animal/cells.

In addition, the peptides of the invention or molecules capable of binding to the peptides may be complexed with toxins, e.g., ricin or cholera, or with other compounds that are toxic to cells. The toxin-binding molecule complex is then targeted to a tumor or other cell by the specificity of the binding molecule for SEQ ID NO: 342-682.

The protein of the invention may also be expressed as a product of transgenic animals, e.g., as a component of the milk of transgenic cows, goats, pigs, or sheep which are characterized by somatic or germ cells containing a nucleotide sequence encoding the protein.

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The proteins provided herein also include proteins characterized by amino acid sequences similar to those of purified proteins but into which modification are naturally provided or deliberately engineered. For example, modifications, in the peptide or DNA sequence, can be made by those skilled in the art using known techniques. Modifications of interest in the protein sequences may include the alteration, substitution, replacement, insertion or deletion of a selected amino acid residue in the coding sequence. For example, one or more of the cysteine residues may be deleted or replaced with another amino acid to alter the conformation of the molecule. Techniques for such alteration, substitution, replacement, insertion or deletion are well known to those skilled in the art (see, e.g., U.S. Pat. No. 4,518,584). Preferably, such alteration, substitution, replacement, insertion or deletion retains the desired activity of the protein. Regions of the protein that are important for the protein function can be determined by various methods known in the art including the alanine-scanning method which involved systematic substitution of single or strings of amino acids with alanine, followed by testing the resulting alanine-containing variant for biological activity. This type of analysis determines the importance of the substituted amino acid(s) in biological activity. Regions of the protein that are important for protein function may be determined by the eMATRIX program.

Other fragments and derivatives of the sequences of proteins which would be expected to retain protein activity in whole or in part and are useful for screening or other immunological methodologies may also be easily made by those skilled in the art given the disclosures herein. Such modifications are encompassed by the present invention.

The protein may also be produced by operably linking the isolated polynucleotide of the invention to suitable control sequences in one or more insect expression vectors, and employing an insect expression system. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, e.g., Invitrogen, San Diego, Calif., U.S.A. (the MaxBat™ kit), and such methods are well known in the art, as described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987), incorporated herein by reference. As used herein, an insect cell capable of expressing a polynucleotide of the present invention is "transformed."

The protein of the invention may be prepared by culturing transformed host cells under culture conditions suitable to express the recombinant protein. The resulting expressed protein may then be purified from such culture (*i.e.*, from culture medium or cell extracts) using known purification processes, such as gel filtration and ion exchange chromatography. The purification of the protein may also include an affinity column containing agents which will bind to the protein; one or more column steps over such affinity resins as concanavalin A-agarose, heparin-toyopearlTM or Cibacrom blue 3GA SepharoseTM; one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether; or immunoaffinity chromatography.

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Alternatively, the protein of the invention may also be expressed in a form which will facilitate purification. For example, it may be expressed as a fusion protein, such as those of maltose binding protein (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX), or as a His tag. Kits for expression and purification of such fusion proteins are commercially available from New England BioLab (Beverly, Mass.), Pharmacia (Piscataway, N.J.) and Invitrogen, respectively. The protein can also be tagged with an epitope and subsequently purified by using a specific antibody directed to such epitope. One such epitope ("FLAG®") is commercially available from Kodak (New Haven, Conn.).

Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify the protein. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous isolated recombinant protein. The protein thus purified is substantially free of other mammalian proteins and is defined in accordance with the present invention as an "isolated protein."

The polypeptides of the invention include analogs (variants). This embraces fragments, as well as peptides in which one or more amino acids has been deleted, inserted, or substituted. Also, analogs of the polypeptides of the invention embrace fusions of the polypeptides or modifications of the polypeptides of the invention, wherein the polypeptide or analog is fused to another moiety or moieties, e.g., targeting moiety or another therapeutic agent. Such analogs may exhibit improved properties such as activity and/or stability. Examples of moieties which may be fused to the polypeptide or an analog include, for example, targeting moieties which provide for the delivery of polypeptide to pancreatic cells, e.g., antibodies to pancreatic cells, antibodies to immune cells such as T-cells, monocytes,

dendritic cells, granulocytes, etc., as well as receptor and ligands expressed on pancreatic or immune cells. Other moieties which may be fused to the polypeptide include therapeutic agents which are used for treatment, for example, immunosuppressive drugs such as cyclosporin, SK506, azathioprine, CD3 antibodies and steroids. Also, polypeptides may be fused to immune modulators, and other cytokines such as alpha or beta interferon.

4.6.1 DETERMINING POLYPEPTIDE AND POLYNUCLEOTIDE IDENTITY AND SIMILARITY

Preferred identity and/or similarity are designed to give the largest match between the 10 sequences tested. Methods to determine identity and similarity are codified in computer programs including, but are not limited to, the GCG program package, including GAP (Devereux, J., et al., Nucleic Acids Research 12(1):387 (1984); Genetics Computer Group. University of Wisconsin, Madison, WI), BLASTP, BLASTN, BLASTX, FASTA (Altschul, S.F. et al., J. Molec. Biol. 215:403-410 (1990), PSI-BLAST (Altschul S.F. et al., Nucleic 15 Acids Res. vol. 25, pp. 3389-3402, herein incorporated by reference), eMatrix software (Wu et al., J. Comp. Biol., Vol. 6, pp. 219-235 (1999), herein incorporated by reference), eMotif software (Nevill-Manning et al, ISMB-97, Vol. 4, pp. 202-209, herein incorporated by reference), pFam software (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1), pp. 320-322 (1998), herein incorporated by reference), the GeneAtlas software (Molecular Simulations 20 Inc. (MSI), San Diego, CA) (Sanchez and Sali (1998) Proc. Natl. Acad. Sci., 95, 13597-13602; Kitson DH et al, (2000) "Remote homology detection using structural modeling - an evaluation" Submitted; Fischer and Eisenberg (1996) Protein Sci. 5, 947-955), Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark), and the Kyte-Doolittle hydrophobocity prediction algorithm (J. Mol Biol, 157, pp. 105-31 (1982), incorporated herein by reference). The 25 BLAST programs are publicly available from the National Center for Biotechnology Information (NCBI) and other sources (BLAST Manual, Altschul, S., et al. NCB NLM NIH Bethesda, MD 20894; Altschul, S., et al., J. Mol. Biol. 215:403-410 (1990).

4.7 CHIMERIC AND FUSION PROTEINS

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The invention also provides chimeric or fusion proteins. As used herein, a "chimeric protein" or "fusion protein" comprises a polypeptide of the invention operatively linked to another polypeptide. Within a fusion protein the polypeptide according to the invention can correspond to all or a portion of a protein according to the invention. In one embodiment, a

fusion protein comprises at least one biologically active portion of a protein according to the invention. In another embodiment, a fusion protein comprises at least two biologically active portions of a protein according to the invention. Within the fusion protein, the term "operatively linked" is intended to indicate that the polypeptide according to the invention and the other polypeptide are fused in-frame to each other. The polypeptide can be fused to the N-terminus or C-terminus.

For example, in one embodiment a fusion protein comprises a polypeptide according to the invention operably linked to the extracellular domain of a second protein. In another embodiment, the fusion protein is a GST-fusion protein in which the polypeptide sequences of the invention are fused to the C-terminus of the GST (i.e., glutathione S-transferase) sequences.

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In another embodiment, the fusion protein is an immunoglobulin fusion protein in which the polypeptide sequences according to the invention comprise one or more domains fused to sequences derived from a member of the immunoglobulin protein family. The immunoglobulin fusion proteins of the invention can be incorporated into pharmaceutical compositions and administered to a subject to inhibit an interaction between a ligand and a protein of the invention on the surface of a cell, to thereby suppress signal transduction in vivo. The immunoglobulin fusion proteins can be used to affect the bioavailability of a cognate ligand. Inhibition of the ligand/protein interaction may be useful therapeutically for both the treatment of proliferative and differentiative disorders, e,g., cancer as well as modulating (e.g., promoting or inhibiting) cell survival. Moreover, the immunoglobulin fusion proteins of the invention can be used as immunogens to produce antibodies in a subject, to purify ligands, and in screening assays to identify molecules that inhibit the interaction of a polypeptide of the invention with a ligand.

A chimeric or fusion protein of the invention can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different polypeptide sequences are ligated together in-frame in accordance with conventional techniques, e.g., by employing blunt-ended or stagger-ended termini for ligation, restriction enzyme digestion to provide for appropriate termini, filling-in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and enzymatic ligation. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers that give rise to complementary overhangs

between two consecutive gene fragments that can subsequently be annealed and reamplified to generate a chimeric gene sequence (see, for example, Ausubel et al. (eds.) CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST polypeptide). A nucleic acid encoding a polypeptide of the invention can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the protein of the invention.

4.8 GENE THERAPY

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Mutations in the polynucleotides of the invention may result in loss of normal function of the encoded protein. The invention thus provides gene therapy to restore normal activity of the polypeptides of the invention; or to treat disease states involving polypeptides of the invention. Delivery of a functional gene encoding polypeptides of the invention to appropriate cells is effected ex vivo, in situ, or in vivo by use of vectors, and more particularly viral vectors (e.g., adenovirus, adeno-associated virus, or a retrovirus), or ex vivo by use of physical DNA transfer methods (e.g., liposomes or chemical treatments). See, for example, Anderson, Nature, supplement to vol. 392, no. 6679, pp.25-20 (1998). For additional reviews of gene therapy technology see Friedmann, Science, 244: 1275-1281 (1989); Verma. Scientific American: 68-84 (1990); and Miller, Nature, 357: 455-460 (1992). Introduction of any one of the nucleotides of the present invention or a gene encoding the polypeptides of the present invention can also be accomplished with extrachromosomal substrates (transient expression) or artificial chromosomes (stable expression). Cells may also be cultured ex vivo in the presence of proteins of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes. Alternatively, it is contemplated that in other human disease states, preventing the expression of or inhibiting the activity of polypeptides of the invention will be useful in treating the disease states. It is contemplated that antisense therapy or gene therapy could be applied to negatively regulate the expression of polypeptides of the invention.

Other methods inhibiting expression of a protein include the introduction of antisense molecules to the nucleic acids of the present invention, their complements, or their translated RNA sequences, by methods known in the art. Further, the polypeptides of the present invention can be inhibited by using targeted deletion methods, or the insertion of a negative regulatory element such as a silencer, which is tissue specific.

The present invention still further provides cells genetically engineered *in vivo* to express the polynucleotides of the invention, wherein such polynucleotides are in operative association with a regulatory sequence heterologous to the host cell which drives expression of the polynucleotides in the cell. These methods can be used to increase or decrease the expression of the polynucleotides of the present invention.

Knowledge of DNA sequences provided by the invention allows for modification of cells to permit, increase, or decrease, expression of endogenous polypeptide. Cells can be modified (e.g., by homologous recombination) to provide increased polypeptide expression by replacing, in whole or in part, the naturally occurring promoter with all or part of a heterologous promoter so that the cells express the protein at higher levels. The heterologous promoter is inserted in such a manner that it is operatively linked to the desired protein encoding sequences. See, for example, PCT International Publication No. WO 94/12650, PCT International Publication No. WO 91/09955. It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (e.g., ada, dhfr, and the multifunctional CAD gene which encodes carbamyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the desired protein coding sequence, amplification of the marker DNA by standard selection methods results in co-amplification of the desired protein coding sequences in the cells.

In another embodiment of the present invention, cells and tissues may be engineered to express an endogenous gene comprising the polynucleotides of the invention under the control of inducible regulatory elements, in which case the regulatory sequences of the endogenous gene may be replaced by homologous recombination. As described herein, gene targeting can be used to replace a gene's existing regulatory region with a regulatory sequence isolated from a different gene or a novel regulatory sequence synthesized by genetic engineering methods. Such regulatory sequences may be comprised of promoters, enhancers, scaffold-attachment regions, negative regulatory elements, transcriptional initiation sites, regulatory protein binding sites or combinations of said sequences. Alternatively, sequences which affect the structure or stability of the RNA or protein produced may be replaced, removed, added, or otherwise modified by targeting. These sequences include polyadenylation signals, mRNA stability elements, splice sites, leader sequences for enhancing or modifying transport or secretion properties of the protein, or other sequences which alter or improve the function or stability of protein or RNA molecules.

The targeting event may be a simple insertion of the regulatory sequence, placing the gene under the control of the new regulatory sequence, e.g., inserting a new promoter or enhancer or both upstream of a gene. Alternatively, the targeting event may be a simple deletion of a regulatory element, such as the deletion of a tissue-specific negative regulatory element. Alternatively, the targeting event may replace an existing element; for example, a tissue-specific enhancer can be replaced by an enhancer that has broader or different cell-type specificity than the naturally occurring elements. Here, the naturally occurring sequences are deleted and new sequences are added. In all cases, the identification of the targeting event may be facilitated by the use of one or more selectable marker genes that are contiguous with the targeting DNA, allowing for the selection of cells in which the exogenous DNA has integrated into the cell genome. The identification of the targeting event may also be facilitated by the use of one or more marker genes exhibiting the property of negative selection, such that the negatively selectable marker is linked to the exogenous DNA, but configured such that the negatively selectable marker flanks the targeting sequence, and such that a correct homologous recombination event with sequences in the host cell genome does not result in the stable integration of the negatively selectable marker. Markers useful for this purpose include the Herpes Simplex Virus thymidine kinase (TK) gene or the bacterial xanthine-guanine phosphoribosyl-transferase (gpt) gene.

The gene targeting or gene activation techniques which can be used in accordance with this aspect of the invention are more particularly described in U.S. Patent No. 5,272,071 to Chappel; U.S. Patent No. 5,578,461 to Sherwin et al.; International Application No. PCT/US92/09627 (WO93/09222) by Selden et al.; and International Application No. PCT/US90/06436 (WO91/06667) by Skoultchi et al., each of which is incorporated by reference herein in its entirety.

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4.9 TRANSGENIC ANIMALS

In preferred methods to determine biological functions of the polypeptides of the invention in vivo, one or more genes provided by the invention are either over expressed or inactivated in the germ line of animals using homologous recombination [Capecchi, Science 244:1288-1292 (1989)]. Animals in which the gene is over expressed, under the regulatory control of exogenous or endogenous promoter elements, are known as transgenic animals. Animals in which an endogenous gene has been inactivated by homologous recombination are referred to as "knockout" animals. Knockout animals, preferably non-human mammals,

can be prepared as described in U.S. Patent No. 5,557,032, incorporated herein by reference. Transgenic animals are useful to determine the roles polypeptides of the invention play in biological processes, and preferably in disease states. Transgenic animals are useful as model systems to identify compounds that modulate lipid metabolism. Transgenic animals, preferably non-human mammals, are produced using methods as described in U.S. Patent No 5,489,743 and PCT Publication No. WO94/28122, incorporated herein by reference.

Transgenic animals can be prepared wherein all or part of a promoter of the polynucleotides of the invention is either activated or inactivated to alter the level of expression of the polypeptides of the invention. Inactivation can be carried out using homologous recombination methods described above. Activation can be achieved by supplementing or even replacing the homologous promoter to provide for increased protein expression. The homologous promoter can be supplemented by insertion of one or more heterologous enhancer elements known to confer promoter activation in a particular tissue.

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The polynucleotides of the present invention also make possible the development, through, e.g., homologous recombination or knock out strategies, of animals that fail to express polypeptides of the invention or that express a variant polypeptide. Such animals are useful as models for studying the *in vivo* activities of polypeptide as well as for studying modulators of the polypeptides of the invention.

In preferred methods to determine biological functions of the polypeptides of the invention *in vivo*, one or more genes provided by the invention are either over expressed or inactivated in the germ line of animals using homologous recombination [Capecchi, Science 244:1288-1292 (1989)]. Animals in which the gene is over expressed, under the regulatory control of exogenous or endogenous promoter elements, are known as transgenic animals. Animals in which an endogenous gene has been inactivated by homologous recombination are referred to as "knockout" animals. Knockout animals, preferably non-human mammals, can be prepared as described in U.S. Patent No. 5,557,032, incorporated herein by reference. Transgenic animals are useful to determine the roles polypeptides of the invention play in biological processes, and preferably in disease states. Transgenic animals are useful as model systems to identify compounds that modulate lipid metabolism. Transgenic animals, preferably non-human mammals, are produced using methods as described in U.S. Patent No 5,489,743 and PCT Publication No. WO94/28122, incorporated herein by reference.

Transgenic animals can be prepared wherein all or part of the polynucleotides of the invention promoter is either activated or inactivated to alter the level of expression of the

polypeptides of the invention. Inactivation can be carried out using homologous recombination methods described above. Activation can be achieved by supplementing or even replacing the homologous promoter to provide for increased protein expression. The homologous promoter can be supplemented by insertion of one or more heterologous enhancer elements known to confer promoter activation in a particular tissue.

4.10 USES AND BIOLOGICAL ACTIVITY

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The polynucleotides and proteins of the present invention are expected to exhibit one or more of the uses or biological activities (including those associated with assays cited herein) identified herein. Uses or activities described for proteins of the present invention may be provided by administration or use of such proteins or of polynucleotides encoding such proteins (such as, for example, in gene therapies or vectors suitable for introduction of DNA). The mechanism underlying the particular condition or pathology will dictate whether the polypeptides of the invention, the polynucleotides of the invention or modulators (activators or inhibitors) thereof would be beneficial to the subject in need of treatment. Thus, "therapeutic compositions of the invention" include compositions comprising isolated polynucleotides (including recombinant DNA molecules, cloned genes and degenerate variants thereof) or polypeptides of the invention (including full length protein, mature protein and truncations or domains thereof), or compounds and other substances that modulate the overall activity of the target gene products, either at the level of target gene/protein expression or target protein activity. Such modulators include polypeptides, analogs, (variants), including fragments and fusion proteins, antibodies and other binding proteins; chemical compounds that directly or indirectly activate or inhibit the polypeptides of the invention (identified, e.g., via drug screening assays as described herein); antisense polynucleotides and polynucleotides suitable for triple helix formation; and in particular antibodies or other binding partners that specifically recognize one or more epitopes of the polypeptides of the invention.

The polypeptides of the present invention may likewise be involved in cellular activation or in one of the other physiological pathways described herein.

4.10.1 RESEARCH USES AND UTILITIES

The polynucleotides provided by the present invention can be used by the research community for various purposes. The polynucleotides can be used to express recombinant

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protein for analysis, characterization or therapeutic use; as markers for tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as molecular weight markers on gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions; to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; as a probe to "subtract-out" known sequences in the process of discovering other novel polynucleotides; for selecting and making oligomers for attachment to a "gene chip" or other support, including for examination of expression patterns; to raise anti-protein antibodies using DNA immunization techniques; and as an antigen to raise anti-DNA antibodies or elicit another immune response. Where the polynucleotide encodes a protein which binds or potentially binds to another protein (such as, for example, in a receptor-ligand interaction), the polynucleotide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803 (1993)) to identify polynucleotides encoding the other protein with which binding occurs or to identify inhibitors of the binding interaction.

The polypeptides provided by the present invention can similarly be used in assays to determine biological activity, including in a panel of multiple proteins for high-throughput screening; to raise antibodies or to elicit another immune response; as a reagent (including the labeled reagent) in assays designed to quantitatively determine levels of the protein (or its receptor) in biological fluids; as markers for tissues in which the corresponding polypeptide is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in a disease state); and, of course, to isolate correlative receptors or ligands. Proteins involved in these binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction.

Any or all of these research utilities are capable of being developed into reagent grade or kit format for commercialization as research products.

Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include without limitation "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E. F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic Press, Berger, S. L. and A. R. Kimmel eds., 1987.

4.10.2 NUTRITIONAL USES

Polynucleotides and polypeptides of the present invention can also be used as nutritional sources or supplements. Such uses include without limitation use as a protein or amino acid supplement, use as a carbon source, use as a nitrogen source and use as a source of carbohydrate. In such cases the polypeptide or polynucleotide of the invention can be added to the feed of a particular organism or can be administered as a separate solid or liquid preparation, such as in the form of powder, pills, solutions, suspensions or capsules. In the case of microorganisms, the polypeptide or polynucleotide of the invention can be added to the medium in or on which the microorganism is cultured.

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4.10.3 CYTOKINE AND CELL PROLIFERATION/DIFFERENTIATION ACTIVITY

A polypeptide of the present invention may exhibit activity relating to cytokine, cell proliferation (either inducing or inhibiting) or cell differentiation (either inducing or inhibiting) activity or may induce production of other cytokines in certain cell populations. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Many protein factors discovered to date, including all known cytokines, have exhibited activity in one or more factor-dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cytokine activity. The activity of therapeutic compositions of the present invention is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+(preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e, CMK, HUVEC, and Caco. Therapeutic compositions of the invention can be used in the following:

Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Bertagnolli et al., J. Immunol. 145:1706-1712, 1990; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Bertagnolli, et al., I. Immunol. 149:3778-3783, 1992; Bowman et al., I. Immunol. 152:1756-1761, 1994.

Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described in: Polyclonal T cell stimulation, Kruisbeek, A. M. and Shevach, E. M. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 3.12.1-3.12.14, John Wiley and Sons, Toronto. 1994; and Measurement of mouse and human interleukin-γ, Schreiber, R. D. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 6.8.1-6.8.8, John Wiley and Sons, Toronto. 1994.

Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described in: Measurement of Human and Murine Interleukin 2 and Interleukin 4, Bottomly, K., Davis, L. S. and Lipsky, P. E. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto. 1991; deVries et al., J. Exp. Med. 173:1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc. Natl. Acad. Sci. U.S.A. 80:2931-2938, 1983; Measurement of mouse and human interleukin 6--Nordan, R. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.6.1-6.6.5, John Wiley and Sons, Toronto. 1991; Smith et al., Proc. Natl. Aced. Sci. U.S.A. 83:1857-1861, 1986; Measurement of human Interleukin 11--Bennett, F., Giannotti, J., Clark, S. C. and Turner, K. J. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto. 1991; Measurement of mouse and human Interleukin 9--Ciarletta, A., Giannotti, J., Clark, S. C. and Turner, K. J. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto. 1991.

Assays for T-cell clone responses to antigens (which will identify, among others, proteins that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and cytokine production) include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function; Chapter 6, Cytokines and their cellular receptors; Chapter 7, Immunologic studies in Humans); Weinberger et al., Proc. Natl. Acad. Sci. USA 77:6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988.

4.10.4 STEM CELL GROWTH FACTOR ACTIVITY

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A polypeptide of the present invention may exhibit stem cell growth factor activity and be involved in the proliferation, differentiation and survival of pluripotent and totipotent

stem cells including primordial germ cells, embryonic stem cells, hematopoietic stem cells and/or germ line stem cells. Administration of the polypeptide of the invention to stem cells in vivo or ex vivo is expected to maintain and expand cell populations in a totipotential or pluripotential state which would be useful for re-engineering damaged or diseased tissues, transplantation, manufacture of bio-pharmaceuticals and the development of bio-sensors. The ability to produce large quantities of human cells has important working applications for the production of human proteins which currently must be obtained from non-human sources or donors, implantation of cells to treat diseases such as Parkinson's, Alzheimer's and other neurodegenerative diseases; tissues for grafting such as bone marrow, skin, cartilage, tendons, bone, muscle (including cardiac muscle), blood vessels, cornea, neural cells, gastrointestinal cells and others; and organs for transplantation such as kidney, liver, pancreas (including islet cells), heart and lung.

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It is contemplated that multiple different exogenous growth factors and/or cytokines may be administered in combination with the polypeptide of the invention to achieve the desired effect, including any of the growth factors listed herein, other stem cell maintenance factors, and specifically including stem cell factor (SCF), leukemia inhibitory factor (LIF), Flt-3 ligand (Flt-3L), any of the interleukins, recombinant soluble IL-6 receptor fused to IL-6, macrophage inflammatory protein 1-alpha (MIP-1-alpha), G-CSF, GM-CSF, thrombopoietin (TPO), platelet factor 4 (PF-4), platelet-derived growth factor (PDGF), neural growth factors and basic fibroblast growth factor (bFGF).

Since totipotent stem cells can give rise to virtually any mature cell type, expansion of these cells in culture will facilitate the production of large quantities of mature cells.

Techniques for culturing stem cells are known in the art and administration of polypeptides of the invention, optionally with other growth factors and/or cytokines, is expected to enhance the survival and proliferation of the stem cell populations. This can be accomplished by direct administration of the polypeptide of the invention to the culture medium.

Alternatively, stroma cells transfected with a polynucleotide that encodes for the polypeptide of the invention can be used as a feeder layer for the stem cell populations in culture or in vivo. Stromal support cells for feeder layers may include embryonic bone marrow fibroblasts, bone marrow stromal cells, fetal liver cells, or cultured embryonic fibroblasts (see U.S. Patent No. 5,690,926).

Stem cells themselves can be transfected with a polynucleotide of the invention to induce autocrine expression of the polypeptide of the invention. This will allow for

generation of undifferentiated totipotential/pluripotential stem cell lines that are useful as is or that can then be differentiated into the desired mature cell types. These stable cell lines can also serve as a source of undifferentiated totipotential/pluripotential mRNA to create cDNA libraries and templates for polymerase chain reaction experiments. These studies would allow for the isolation and identification of differentially expressed genes in stem cell populations that regulate stem cell proliferation and/or maintenance.

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Expansion and maintenance of totipotent stem cell populations will be useful in the treatment of many pathological conditions. For example, polypeptides of the present invention may be used to manipulate stem cells in culture to give rise to neuroepithelial cells that can be used to augment or replace cells damaged by illness, autoimmune disease, accidental damage or genetic disorders. The polypeptide of the invention may be useful for inducing the proliferation of neural cells and for the regeneration of nerve and brain tissue, i.e. for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders which involve degeneration, death or trauma to neural cells or nerve tissue. In addition, the expanded stem cell populations can also be genetically altered for gene therapy purposes and to decrease host rejection of replacement tissues after grafting or implantation.

Expression of the polypeptide of the invention and its effect on stem cells can also be manipulated to achieve controlled differentiation of the stem cells into more differentiated cell types. A broadly applicable method of obtaining pure populations of a specific differentiated cell type from undifferentiated stem cell populations involves the use of a cell-type specific promoter driving a selectable marker. The selectable marker allows only cells of the desired type to survive. For example, stem cells can be induced to differentiate into cardiomyocytes (Wobus et al., Differentiation, 48: 173-182, (1991); Klug et al., J. Clin. Invest., 98(1): 216-224, (1998)) or skeletal muscle cells (Browder, L. W. In: *Principles of Tissue Engineering eds.* Lanza et al., Academic Press (1997)). Alternatively, directed differentiation of stem cells can be accomplished by culturing the stem cells in the presence of a differentiation factor such as retinoic acid and an antagonist of the polypeptide of the invention which would inhibit the effects of endogenous stem cell factor activity and allow differentiation to proceed.

In vitro cultures of stem cells can be used to determine if the polypeptide of the invention exhibits stem cell growth factor activity. Stem cells are isolated from any one of various cell sources (including hematopoietic stem cells and embryonic stem cells) and

cultured on a feeder layer, as described by Thompson et al. Proc. Natl. Acad. Sci, U.S.A., 92: 7844-7848 (1995), in the presence of the polypeptide of the invention alone or in combination with other growth factors or cytokines. The ability of the polypeptide of the invention to induce stem cells proliferation is determined by colony formation on semi-solid support e.g. as described by Bernstein et al., Blood, 77: 2316-2321 (1991).

4.10.5 HEMATOPOIESIS REGULATING ACTIVITY

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A polypeptide of the present invention may be involved in regulation of hematopoiesis and, consequently, in the treatment of myeloid or lymphoid cell disorders. Even marginal biological activity in support of colony forming cells or of factor-dependent cell lines indicates involvement in regulating hematopoiesis, e.g. in supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other cytokines. thereby indicating utility, for example, in treating various anemias or for use in conjunction with irradiation/chemotherapy to stimulate the production of erythroid precursors and/or erythroid cells; in supporting the growth and proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e., traditional CSF activity) useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelo-suppression; in supporting the growth and proliferation of megakaryocytes and consequently of platelets thereby allowing prevention or treatment of various platelet disorders such as thrombocytopenia, and generally for use in place of or complimentary to platelet transfusions; and/or in supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned hematopoietic cells and therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantation, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post irradiation/chemotherapy, either in-vivo or ex-vivo (i.e., in conjunction with bone marrow transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy.

Therapeutic compositions of the invention can be used in the following:

Suitable assays for proliferation and differentiation of various hematopoietic lines are cited above.

Assays for embryonic stem cell differentiation (which will identify, among others, proteins that influence embryonic differentiation hematopoiesis) include, without limitation,

those described in: Johansson et al. Cellular Biology 15:141-151, 1995; Keller et al., Molecular and Cellular Biology 13:473-486, 1993; McClanahan et al., Blood 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, proteins that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, M. G. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 265-268, Wiley-Liss, Inc., New York, N.Y. 1994; Hirayama et al., Proc. Natl. Acad. Sci. USA 89:5907-5911, 1992; Primitive hematopoietic colony forming cells with high proliferative potential, McNiece, I. K. and Briddell, R. A. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 23-39, Wiley-Liss, Inc., New York, N.Y. 1994; Neben et al., Experimental Hematology 22:353-359, 1994; Cobblestone area forming cell assay, Ploemacher, R. E. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 1-21, Wiley-Liss, Inc., New York, N.Y. 1994; Long term bone marrow cultures in the presence of stromal cells, Spooncer, E., Dexter, M. and Allen, T. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 163-179, Wiley-Liss, Inc., New York, N.Y. 1994; Long term culture initiating cell assay, Sutherland, H. J. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 139-162, Wiley-Liss, Inc., New York, N.Y. 1994.

20 4.10.6 TISSUE GROWTH ACTIVITY

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A polypeptide of the present invention also may be involved in bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as in wound healing and tissue repair and replacement, and in healing of burns, incisions and ulcers.

A polypeptide of the present invention which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Compositions of a polypeptide, antibody, binding partner, or other modulator of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery.

A polypeptide of this invention may also be involved in attracting bone-forming cells, stimulating growth of bone-forming cells, or inducing differentiation of progenitors of

bone-forming cells. Treatment of osteoporosis, osteoarthritis, bone degenerative disorders, or periodontal disease, such as through stimulation of bone and/or cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes may also be possible using the composition of the invention.

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Another category of tissue regeneration activity that may involve the polypeptide of the present invention is tendon/ligament formation. Induction of tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue, as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue formation induced by a composition of the present invention contributes to the repair of congenital, trauma induced, or other tendon or ligament defects of other origin, and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments. The compositions of the present invention may provide environment to attract tendon- or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors ex vivo for return in vivo to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendinitis, carpal tunnel syndrome and other tendon or ligament defects. The compositions may also include an appropriate matrix and/or sequestering agent as a carrier as is well known in the art.

The compositions of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve and brain tissue, i.e. for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a composition may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nerve injuries, peripheral neuropathy and localized neuropathies, and central nervous system diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular diseases such as

stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a composition of the invention.

Compositions of the invention may also be useful to promote better or faster closure of non-healing wounds, including without limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

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Compositions of the present invention may also be involved in the generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, intestine, kidney, skin, endothelium), muscle (smooth, skeletal or cardiac) and vascular (including vascular endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part of the desired effects may be by inhibition or modulation of fibrotic scarring may allow normal tissue to regenerate. A polypeptide of the present invention may also exhibit angiogenic activity.

A composition of the present invention may also be useful for gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokine damage.

A composition of the present invention may also be useful for promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

Therapeutic compositions of the invention can be used in the following:

Assays for tissue generation activity include, without limitation, those described in: International Patent Publication No. WO95/16035 (bone, cartilage, tendon); International Patent Publication No. WO95/05846 (nerve, neuronal); International Patent Publication No. WO91/07491 (skin, endothelium).

Assays for wound healing activity include, without limitation, those described in: Winter, Epidermal Wound Healing, pps. 71-112 (Maibach, H. I. and Rovee, D. T., eds.), Year Book Medical Publishers, Inc., Chicago, as modified by Eaglstein and Mertz, J. Invest. Dermatol 71:382-84 (1978).

4.10.7 IMMUNE STIMULATING OR SUPPRESSING ACTIVITY

A polypeptide of the present invention may also exhibit immune stimulating or immune suppressing activity, including without limitation the activities for which assays are described herein. A polynucleotide of the invention can encode a polypeptide exhibiting such activities. A protein may be useful in the treatment of various immune deficiencies and

disorders (including severe combined immunodeficiency (SCID)), e.g., in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by viral (e.g., HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious diseases causes by viral, bacterial, fungal or other infection may be treatable using a protein of the present invention, including infections by HIV, hepatitis viruses, herpes viruses, mycobacteria, Leishmania spp., malaria spp. and various fungal infections such as candidiasis. Of course, in this regard, proteins of the present invention may also be useful where a boost to the immune system generally may be desirable, i.e., in the treatment of cancer.

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Autoimmune disorders which may be treated using a protein of the present invention include, for example, connective tissue disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitis, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a protein (or antagonists thereof, including antibodies) of the present invention may also to be useful in the treatment of allergic reactions and conditions (e.g., anaphylaxis, serum sickness, drug reactions, food allergies, insect venom allergies, mastocytosis, allergic rhinitis, hypersensitivity pneumonitis, urticaria, angioedema, eczema, atopic dermatitis, allergic contact dermatitis, erythema multiforme, Stevens-Johnson syndrome, allergic conjunctivitis, atopic keratoconjunctivitis, venereal keratoconjunctivitis, giant papillary conjunctivitis and contact allergies), such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired (including, for example, organ transplantation), may also be treatable using a protein (or antagonists thereof) of the present invention. The therapeutic effects of the polypeptides or antagonists thereof on allergic reactions can be evaluated by in vivo animals models such as the cumulative contact enhancement test (Lastbom et al., Toxicology 125: 59-66, 1998), skin prick test (Hoffmann et al., Allergy 54: 446-54, 1999), guinea pig skin sensitization test (Vohr et al., Arch. Toxocol. 73: 501-9), and murine local lymph node assay (Kimber et al., J. Toxicol. Environ. Health 53: 563-79).

Using the proteins of the invention it may also be possible to modulate immune responses, in a number of ways. Down regulation may be in the form of inhibiting or blocking an immune response already in progress or may involve preventing the induction of

an immune response. The functions of activated T cells may be inhibited by suppressing T cell responses or by inducing specific tolerance in T cells, or both. Immunosuppression of T cell responses is generally an active, non-antigen-specific, process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or anergy in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after exposure to the tolerizing agent has ceased. Operationally, tolerance can be demonstrated by the lack of a T cell response upon reexposure to specific antigen in the absence of the tolerizing agent.

Down regulating or preventing one or more antigen functions (including without limitation B lymphocyte antigen functions (such as, for example, B7)), e.g., preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a therapeutic composition of the invention may prevent cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, a lack of costimulation may also be sufficient to anergize the T cells, thereby inducing tolerance in a subject. Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of a combination of B lymphocyte antigens.

The efficacy of particular therapeutic compositions in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the immunosuppressive effects of CTLA4Ig fusion proteins in vivo as described in Lenschow et al., Science 257:789-792 (1992) and Turka et al., Proc. Natl. Acad. Sci USA, 89:11102-11105 (1992). In addition, murine models of GVHD (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of therapeutic compositions of the invention on the development of that disease.

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Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells that are reactive against self tissue and which promote the production of cytokines and autoantibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms. Administration of reagents which block stimulation of T cells can be used to inhibit T cell activation and prevent production of autoantibodies or T cell-derived cytokines which may be involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases. Examples include murine experimental autoimmune encephalitis, systemic lupus erythmatosis in MRL/lpr/lpr mice or NZB hybrid mice, murine autoimmune collagen arthritis, diabetes mellitus in NOD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 840-856).

Upregulation of an antigen function (e.g., a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune responses may be in the form of enhancing an existing immune response or eliciting an initial immune response. For example, enhancing an immune response may be useful in cases of viral infection, including systemic viral diseases such as influenza, the common cold, and encephalitis.

Alternatively, anti-viral immune responses may be enhanced in an infected patient by removing T cells from the patient, costimulating the T cells in vitro with viral antigen-pulsed APCs either expressing a peptide of the present invention or together with a stimulatory form of a soluble peptide of the present invention and reintroducing the in vitro activated T cells into the patient. Another method of enhancing anti-viral immune responses would be to isolate infected cells from a patient, transfect them with a nucleic acid encoding a protein of the present invention as described herein such that the cells express all or a portion of the protein on their surface, and reintroduce the transfected cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to, and thereby activate, T cells in vivo.

A polypeptide of the present invention may provide the necessary stimulation signal to T cells to induce a T cell mediated immune response against the transfected tumor cells. In

addition, tumor cells which lack MHC class I or MHC class II molecules, or which fail to reexpress sufficient mounts of MHC class I or MHC class II molecules, can be transfected with nucleic acid encoding all or a portion of (e.g., a cytoplasmic-domain truncated portion) of an MHC class I alpha chain protein and β₂ microglobulin protein or an MHC class II alpha chain protein and an MHC class II beta chain protein to thereby express MHC class I or MHC class II proteins on the cell surface. Expression of the appropriate class I or class II MHC in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor associated antigens and induce tumor specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumor-specific tolerance in the subject.

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The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., I. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bowman et al., J. Virology 61:1992-1998; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, proteins that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, J. Immunol. 144:3028-3033, 1990; and Assays for B cell function: In vitro antibody production, Mond, J. J. and Brunswick, M. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto. 1994.

Mixed lymphocyte reaction (MLR) assays (which will identify, among others, proteins that generate predominantly Th1 and CTL responses) include, without limitation,

those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, proteins expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J. Immunol. 134:536-544, 1995; Inaba et al., Journal of Experimental Medicine 173:549-559, 1991; Macatonia et al., Journal of Immunology 154:5071-5079, 1995; Porgador et al., Journal of Experimental Medicine 182:255-260, 1995; Nair et al., Journal of Virology 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., Journal of Experimental Medicine 169:1255-1264, 1989; Bhardwaj et al., Journal of Clinical Investigation 94:797-807, 1994; and Inaba et al., Journal of Experimental Medicine 172:631-640, 1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, proteins that prevent apoptosis after superantigen induction and proteins that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Research 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, Journal of Immunology 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., International Journal of Oncology 1:639-648, 1992.

Assays for proteins that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cellular Immunology 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc. Nat. Acad Sci. USA 88:7548-7551, 1991.

4.10.8 ACTIVIN/INHIBIN ACTIVITY

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A polypeptide of the present invention may also exhibit activin- or inhibin-related activities. A polynucleotide of the invention may encode a polypeptide exhibiting such characteristics. Inhibins are characterized by their ability to inhibit the release of follicle stimulating hormone (FSH), while activins and are characterized by their ability to stimulate the release of follicle stimulating hormone (FSH). Thus, a polypeptide of the present

invention, alone or in heterodimers with a member of the inhibin family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the polypeptide of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, U.S. Pat. No. 4,798,885. A polypeptide of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as, but not limited to, cows, sheep and pigs.

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The activity of a polypeptide of the invention may, among other means, be measured by the following methods.

Assays for activin/inhibin activity include, without limitation, those described in: Vale et al., Endocrinology 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc. Natl. Acad. Sci. USA 83:3091-3095, 1986.

4.10.9 CHEMOTACTIC/CHEMOKINETIC ACTIVITY

A polypeptide of the present invention may be involved in chemotactic or chemokinetic activity for mammalian cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Chemotactic and chemokinetic receptor activation can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic compositions (e.g. proteins, antibodies, binding partners, or modulators of the invention) provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example, attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of

cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

Therapeutic compositions of the invention can be used in the following:

Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Marguiles, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 6.12, Measurement of alpha and beta Chemokines 6.12.1-6.12.28; Taub et al. J. Clin. Invest. 95:1370-1376, 1995; Lind et al. APMIS 103:140-146, 1995; Muller et al Eur. J. Immunol. 25:1744-1748; Gruber et al. J. of Immunol. 152:5860-5867, 1994; Johnston et al. J. of Immunol. 153:1762-1768, 1994.

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4.10.10 HEMOSTATIC AND THROMBOLYTIC ACTIVITY

A polypeptide of the invention may also be involved in hemostasis or thrombolysis or thrombosis. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Compositions may be useful in treatment of various coagulation disorders (including hereditary disorders, such as hemophilias) or to enhance coagulation and other hemostatic events in treating wounds resulting from trauma, surgery or other causes. A composition of the invention may also be useful for dissolving or inhibiting formation of thromboses and for treatment and prevention of conditions resulting therefrom (such as, for example, infarction of cardiac and central nervous system vessels (e.g., stroke).

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Therapeutic compositions of the invention can be used in the following:

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res. 45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

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4.10.11 CANCER DIAGNOSIS AND THERAPY

Polypeptides of the invention may be involved in cancer cell generation, proliferation or metastasis. Detection of the presence or amount of polynucleotides or polypeptides of the

invention may be useful for the diagnosis and/or prognosis of one or more types of cancer. For example, the presence or increased expression of a polynucleotide/polypeptide of the invention may indicate a hereditary risk of cancer, a precancerous condition, or an ongoing malignancy. Conversely, a defect in the gene or absence of the polypeptide may be associated with a cancer condition. Identification of single nucleotide polymorphisms associated with cancer or a predisposition to cancer may also be useful for diagnosis or prognosis.

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Cancer treatments promote tumor regression by inhibiting tumor cell proliferation, inhibiting angiogenesis (growth of new blood vessels that is necessary to support tumor growth) and/or prohibiting metastasis by reducing tumor cell motility or invasiveness. Therapeutic compositions of the invention may be effective in adult and pediatric oncology including in solid phase tumors/malignancies, locally advanced tumors, human soft tissue sarcomas, metastatic cancer, including lymphatic metastases, blood cell malignancies including multiple myeloma, acute and chronic leukemias, and lymphomas, head and neck cancers including mouth cancer, larynx cancer and thyroid cancer, lung cancers including small cell carcinoma and non-small cell cancers, breast cancers including small cell carcinoma and ductal carcinoma, gastrointestinal cancers including esophageal cancer, stomach cancer, colon cancer, colorectal cancer and polyps associated with colorectal neoplasia, pancreatic cancers, liver cancer, urologic cancers including bladder cancer and prostate cancer, malignancies of the female genital tract including ovarian carcinoma, uterine (including endometrial) cancers, and solid tumor in the ovarian follicle, kidney cancers including renal cell carcinoma, brain cancers including intrinsic brain tumors, neuroblastoma, astrocytic brain tumors, gliomas, metastatic tumor cell invasion in the central nervous system, bone cancers including osteomas, skin cancers including malignant melanoma, tumor progression of human skin keratinocytes, squamous cell carcinoma, basal cell carcinoma, hemangiopericytoma and Karposi's sarcoma.

Polypeptides, polynucleotides, or modulators of polypeptides of the invention (including inhibitors and stimulators of the biological activity of the polypeptide of the invention) may be administered to treat cancer. Therapeutic compositions can be administered in therapeutically effective dosages alone or in combination with adjuvant cancer therapy such as surgery, chemotherapy, radiotherapy, thermotherapy, and laser therapy, and may provide a beneficial effect, e.g. reducing tumor size, slowing rate of tumor growth, inhibiting metastasis, or otherwise improving overall clinical condition, without

necessarily eradicating the cancer.

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The composition can also be administered in therapeutically effective amounts as a portion of an anti-cancer cocktail. An anti-cancer cocktail is a mixture of the polypeptide or modulator of the invention with one or more anti-cancer drugs in addition to a pharmaceutically acceptable carrier for delivery. The use of anti-cancer cocktails as a cancer treatment is routine. Anti-cancer drugs that are well known in the art and can be used as a treatment in combination with the polypeptide or modulator of the invention include: Actinomycin D, Aminoglutethimide, Asparaginase, Bleomycin, Busulfan, Carboplatin, Carmustine, Chlorambucil, Cisplatin (cis-DDP), Cyclophosphamide, Cytarabine HCl (Cytosine arabinoside), Dacarbazine, Dactinomycin, Daunorubicin HCl, Doxorubicin HCl, Estramustine phosphate sodium, Etoposide (V16-213), Floxuridine, 5-Fluorouracil (5-Fu), Flutamide, Hydroxyurea (hydroxycarbamide), Ifosfamide, Interferon Alpha-2a, Interferon Alpha-2b, Leuprolide acetate (LHRH-releasing factor analog), Lomustine, Mechlorethamine HCl (nitrogen mustard), Melphalan, Mercaptopurine, Mesna, Methotrexate (MTX), Mitomycin, Mitoxantrone HCl, Octreotide, Plicamycin, Procarbazine HCl, Streptozocin, Tamoxifen citrate, Thioguanine, Thiotepa, Vinblastine sulfate, Vincristine sulfate, Amsacrine, Azacitidine, Hexamethylmelamine, Interleukin-2, Mitoguazone, Pentostatin, Semustine, Teniposide, and Vindesine sulfate.

In addition, therapeutic compositions of the invention may be used for prophylactic treatment of cancer. There are hereditary conditions and/or environmental situations (e.g. exposure to carcinogens) known in the art that predispose an individual to developing cancers. Under these circumstances, it may be beneficial to treat these individuals with therapeutically effective doses of the polypeptide of the invention to reduce the risk of developing cancers.

In vitro models can be used to determine the effective doses of the polypeptide of the invention as a potential cancer treatment. These in vitro models include proliferation assays of cultured tumor cells, growth of cultured tumor cells in soft agar (see Freshney, (1987) Culture of Animal Cells: A Manual of Basic Technique, Wily-Liss, New York, NY Ch 18 and Ch 21), tumor systems in nude mice as described in Giovanella et al., J. Natl. Can. Inst., 52: 921-30 (1974), mobility and invasive potential of tumor cells in Boyden Chamber assays as described in Pilkington et al., Anticancer Res., 17: 4107-9 (1997), and angiogenesis assays such as induction of vascularization of the chick chorioallantoic membrane or induction of vascular endothelial cell migration as described in Ribatta et al., Intl. J. Dev. Biol., 40: 1189-

97 (1999) and Li et al., Clin. Exp. Metastasis, 17:423-9 (1999), respectively. Suitable tumor cells lines are available, e.g. from American Type Tissue Culture Collection catalogs.

4.10.12 RECEPTOR/LIGAND ACTIVITY

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A polypeptide of the present invention may also demonstrate activity as receptor, receptor ligand or inhibitor or agonist of receptor/ligand interactions. A polynucleotide of the invention can encode a polypeptide exhibiting such characteristics. Examples of such receptors and ligands include, without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell-cell interactions and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses. Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions.

The activity of a polypeptide of the invention may, among other means, be measured by the following methods:

Suitable assays for receptor-ligand activity include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of Cellular Adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J. Immunol. Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

By way of example, the polypeptides of the invention may be used as a receptor for a ligand(s) thereby transmitting the biological activity of that ligand(s). Ligands may be identified through binding assays, affinity chromatography, dihybrid screening assays, BIAcore assays, gel overlay assays, or other methods known in the art.

Studies characterizing drugs or proteins as agonist or antagonist or partial agonists or a partial antagonist require the use of other proteins as competing ligands. The polypeptides of the present invention or ligand(s) thereof may be labeled by being coupled to radioisotopes, colorimetric molecules or a toxin molecules by conventional methods. ("Guide

to Protein Purification" Murray P. Deutscher (ed) Methods in Enzymology Vol. 182 (1990) Academic Press, Inc. San Diego). Examples of radioisotopes include, but are not limited to, tritium and carbon-14. Examples of colorimetric molecules include, but are not limited to, fluorescent molecules such as fluorescamine, or rhodamine or other colorimetric molecules. Examples of toxins include, but are not limited, to ricin.

4.10.13 DRUG SCREENING

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This invention is particularly useful for screening chemical compounds by using the novel polypeptides or binding fragments thereof in any of a variety of drug screening techniques. The polypeptides or fragments employed in such a test may either be free in solution, affixed to a solid support, borne on a cell surface or located intracellularly. One method of drug screening utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant nucleic acids expressing the polypeptide or a fragment thereof. Drugs are screened against such transformed cells in competitive binding assays. Such cells, either in viable or fixed form, can be used for standard binding assays. One may measure, for example, the formation of complexes between polypeptides of the invention or fragments and the agent being tested or examine the diminution in complex formation between the novel polypeptides and an appropriate cell line, which are well known in the art.

Sources for test compounds that may be screened for ability to bind to or modulate (i.e., increase or decrease) the activity of polypeptides of the invention include (1) inorganic and organic chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of either random or mimetic peptides, oligonucleotides or organic molecules.

Chemical libraries may be readily synthesized or purchased from a number of commercial sources, and may include structural analogs of known compounds or compounds that are identified as "hits" or "leads" via natural product screening.

The sources of natural product libraries are microorganisms (including bacteria and fungi), animals, plants or other vegetation, or marine organisms, and libraries of mixtures for screening may be created by: (1) fermentation and extraction of broths from soil, plant or marine microorganisms or (2) extraction of the organisms themselves. Natural product libraries include polyketides, non-ribosomal peptides, and (non-naturally occurring) variants thereof. For a review, see *Science* 282:63-68 (1998).

Combinatorial libraries are composed of large numbers of peptides, oligonucleotides or organic compounds and can be readily prepared by traditional automated synthesis

methods, PCR, cloning or proprietary synthetic methods. Of particular interest are peptide and oligonucleotide combinatorial libraries. Still other libraries of interest include peptide, protein, peptidomimetic, multiparallel synthetic collection, recombinatorial, and polypeptide libraries. For a review of combinatorial chemistry and libraries created therefrom, see Myers, Curr. Opin. Biotechnol. 8:701-707 (1997). For reviews and examples of peptidomimetic libraries, see Al-Obeidi et al., Mol. Biotechnol, 9(3):205-23 (1998); Hruby et al., Curr Opin Chem Biol, 1(1):114-19 (1997); Dorner et al., Bioorg Med Chem, 4(5):709-15 (1996) (alkylated dipeptides).

Identification of modulators through use of the various libraries described herein permits modification of the candidate "hit" (or "lead") to optimize the capacity of the "hit" to bind a polypeptide of the invention. The molecules identified in the binding assay are then tested for antagonist or agonist activity in *in vivo* tissue culture or animal models that are well known in the art. In brief, the molecules are titrated into a plurality of cell cultures or animals and then tested for either cell/animal death or prolonged survival of the animal/cells.

The binding molecules thus identified may be complexed with toxins, e.g., ricin or cholera, or with other compounds that are toxic to cells such as radioisotopes. The toxin-binding molecule complex is then targeted to a tumor or other cell by the specificity of the binding molecule for a polypeptide of the invention. Alternatively, the binding molecules may be complexed with imaging agents for targeting and imaging purposes.

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4.10.14 ASSAY FOR RECEPTOR ACTIVITY

The invention also provides methods to detect specific binding of a polypeptide e.g. a ligand or a receptor. The art provides numerous assays particularly useful for identifying previously unknown binding partners for receptor polypeptides of the invention. For example, expression cloning using mammalian or bacterial cells, or dihybrid screening assays can be used to identify polynucleotides encoding binding partners. As another example, affinity chromatography with the appropriate immobilized polypeptide of the invention can be used to isolate polypeptides that recognize and bind polypeptides of the invention. There are a number of different libraries used for the identification of compounds, and in particular small molecules, that modulate (*i.e.*, increase or decrease) biological activity of a polypeptide of the invention. Ligands for receptor polypeptides of the invention can also be identified by adding exogenous ligands, or cocktails of ligands to two cells populations that are genetically identical except for the expression of the receptor of the invention: one cell population

expresses the receptor of the invention whereas the other does not. The response of the two cell populations to the addition of ligands(s) are then compared. Alternatively, an expression library can be co-expressed with the polypeptide of the invention in cells and assayed for an autocrine response to identify potential ligand(s). As still another example, BIAcore assays, gel overlay assays, or other methods known in the art can be used to identify binding partner polypeptides, including, (1) organic and inorganic chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of random peptides, oligonucleotides or organic molecules.

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The role of downstream intracellular signaling molecules in the signaling cascade of the polypeptide of the invention can be determined. For example, a chimeric protein in which the cytoplasmic domain of the polypeptide of the invention is fused to the extracellular portion of a protein, whose ligand has been identified, is produced in a host cell. The cell is then incubated with the ligand specific for the extracellular portion of the chimeric protein, thereby activating the chimeric receptor. Known downstream proteins involved in intracellular signaling can then be assayed for expected modifications i.e. phosphorylation. Other methods known to those in the art can also be used to identify signaling molecules involved in receptor activity.

4.10.15 ANTI-INFLAMMATORY ACTIVITY

Compositions of the present invention may also exhibit anti-inflammatory activity. The anti-inflammatory activity may be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell-cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Compositions with such activities can be used to treat inflammatory conditions including chronic or acute conditions), including without limitation intimation associated with infection (such as septic shock, sepsis or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of cytokines such as TNF or IL-1. Compositions of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material. Compositions of this

invention may be utilized to prevent or treat conditions such as, but not limited to, sepsis, acute pancreatitis, endotoxin shock, cytokine induced shock, rheumatoid arthritis, chronic inflammatory arthritis, pancreatic cell damage from diabetes mellitus type 1, graft versus host disease, inflammatory bowel disease, inflamation associated with pulmonary disease, other autoimmune disease or inflammatory disease, an antiproliferative agent such as for acute or chronic mylegenous leukemia or in the prevention of premature labor secondary to intrauterine infections.

4.10.16 LEUKEMIAS

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Leukemias and related disorders may be treated or prevented by administration of a therapeutic that promotes or inhibits function of the polynucleotides and/or polypeptides of the invention. Such leukemias and related disorders include but are not limited to acute leukemia, acute lymphocytic leukemia, acute myelocytic leukemia, myeloblastic, promyelocytic, myelomonocytic, monocytic, erythroleukemia, chronic leukemia, chronic myelocytic (granulocytic) leukemia and chronic lymphocytic leukemia (for a review of such disorders, see Fishman et al., 1985, Medicine, 2d Ed., J.B. Lippincott Co., Philadelphia).

4.10.17 NERVOUS SYSTEM DISORDERS

Nervous system disorders, involving cell types which can be tested for efficacy of intervention with compounds that modulate the activity of the polynucleotides and/or polypeptides of the invention, and which can be treated upon thus observing an indication of therapeutic utility, include but are not limited to nervous system injuries, and diseases or disorders which result in either a disconnection of axons, a diminution or degeneration of neurons, or demyelination. Nervous system lesions which may be treated in a patient (including human and non-human mammalian patients) according to the invention include but are not limited to the following lesions of either the central (including spinal cord, brain) or peripheral nervous systems:

- (i) traumatic lesions, including lesions caused by physical injury or associated with surgery, for example, lesions which sever a portion of the nervous system, or compression injuries;
- (ii) ischemic lesions, in which a lack of oxygen in a portion of the nervous system results in neuronal injury or death, including cerebral infarction or ischemia, or spinal cord infarction or ischemia;

(iii) infectious lesions, in which a portion of the nervous system is destroyed or injured as a result of infection, for example, by an abscess or associated with infection by human immunodeficiency virus, herpes zoster, or herpes simplex virus or with Lyme disease, tuberculosis, syphilis;

(iv) degenerative lesions, in which a portion of the nervous system is destroyed or injured as a result of a degenerative process including but not limited to degeneration associated with Parkinson's disease, Alzheimer's disease, Huntington's chorea, or amyotrophic lateral sclerosis;

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- (v) lesions associated with nutritional diseases or disorders, in which a portion of the nervous system is destroyed or injured by a nutritional disorder or disorder of metabolism including but not limited to, vitamin B12 deficiency, folic acid deficiency, Wernicke disease, tobacco-alcohol amblyopia, Marchiafava-Bignami disease (primary degeneration of the corpus callosum), and alcoholic cerebellar degeneration;
- (vi) neurological lesions associated with systemic diseases including but not limited to diabetes (diabetic neuropathy, Bell's palsy), systemic lupus erythematosus, carcinoma, or sarcoidosis;
- (vii) lesions caused by toxic substances including alcohol, lead, or particular neurotoxins; and
- (viii) demyelinated lesions in which a portion of the nervous system is destroyed or injured by a demyelinating disease including but not limited to multiple sclerosis, human immunodeficiency virus-associated myelopathy, transverse myelopathy or various etiologies, progressive multifocal leukoencephalopathy, and central pontine myelinolysis.

Therapeutics which are useful according to the invention for treatment of a nervous system disorder may be selected by testing for biological activity in promoting the survival or differentiation of neurons. For example, and not by way of limitation, therapeutics which elicit any of the following effects may be useful according to the invention:

- (i) increased survival time of neurons in culture;
- (ii) increased sprouting of neurons in culture or in vivo;
- (iii) increased production of a neuron-associated molecule in culture or *in vivo*, e.g., choline acetyltransferase or acetylcholinesterase with respect to motor neurons; or
 - (iv) decreased symptoms of neuron dysfunction in vivo.

Such effects may be measured by any method known in the art. In preferred, non-limiting embodiments, increased survival of neurons may be measured by the method set

forth in Arakawa et al. (1990, J. Neurosci. 10:3507-3515); increased sprouting of neurons may be detected by methods set forth in Pestronk et al. (1980, Exp. Neurol. 70:65-82) or Brown et al. (1981, Ann. Rev. Neurosci. 4:17-42); increased production of neuron-associated molecules may be measured by bioassay, enzymatic assay, antibody binding, Northern blot assay, etc., depending on the molecule to be measured; and motor neuron dysfunction may be measured by assessing the physical manifestation of motor neuron disorder, e.g., weakness, motor neuron conduction velocity, or functional disability.

In specific embodiments, motor neuron disorders that may be treated according to the invention include but are not limited to disorders such as infarction, infection, exposure to toxin, trauma, surgical damage, degenerative disease or malignancy that may affect motor neurons as well as other components of the nervous system, as well as disorders that selectively affect neurons such as amyotrophic lateral sclerosis, and including but not limited to progressive spinal muscular atrophy, progressive bulbar palsy, primary lateral sclerosis, infantile and juvenile muscular atrophy, progressive bulbar paralysis of childhood (Fazio-Londe syndrome), poliomyelitis and the post polio syndrome, and Hereditary Motorsensory Neuropathy (Charcot-Marie-Tooth Disease).

4.10.18 OTHER ACTIVITIES

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A polypeptide of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents, including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in bone form or shape); effecting biorhythms or circadian cycles or rhythms; effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, co-factors or other nutritional factors or component(s); effecting behavioral characteristics, including, without limitation, appetite, libido, stress, cognition (including cognitive disorders), depression (including depressive disorders) and violent behaviors; providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting deficiencies of

the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein.

4.10.19 IDENTIFICATION OF POLYMORPHISMS

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The demonstration of polymorphisms makes possible the identification of such polymorphisms in human subjects and the pharmacogenetic use of this information for diagnosis and treatment. Such polymorphisms may be associated with, e.g., differential predisposition or susceptibility to various disease states (such as disorders involving inflammation or immune response) or a differential response to drug administration, and this genetic information can be used to tailor preventive or therapeutic treatment appropriately. For example, the existence of a polymorphism associated with a predisposition to inflammation or autoimmune disease makes possible the diagnosis of this condition in humans by identifying the presence of the polymorphism.

Polymorphisms can be identified in a variety of ways known in the art which all generally involve obtaining a sample from a patient, analyzing DNA from the sample. optionally involving isolation or amplification of the DNA, and identifying the presence of the polymorphism in the DNA. For example, PCR may be used to amplify an appropriate fragment of genomic DNA which may then be sequenced. Alternatively, the DNA may be subjected to allele-specific oligonucleotide hybridization (in which appropriate oligonucleotides are hybridized to the DNA under conditions permitting detection of a single base mismatch) or to a single nucleotide extension assay (in which an oligonucleotide that hybridizes immediately adjacent to the position of the polymorphism is extended with one or more labeled nucleotides). In addition, traditional restriction fragment length polymorphism analysis (using restriction enzymes that provide differential digestion of the genomic DNA depending on the presence or absence of the polymorphism) may be performed. Arrays with nucleotide sequences of the present invention can be used to detect polymorphisms. The array can comprise modified nucleotide sequences of the present invention in order to detect the nucleotide sequences of the present invention. In the alternative, any one of the nucleotide sequences of the present invention can be placed on the array to detect changes from those sequences.

Alternatively a polymorphism resulting in a change in the amino acid sequence could also be detected by detecting a corresponding change in amino acid sequence of the protein, e.g., by an antibody specific to the variant sequence.

4.10.20 ARTHRITIS AND INFLAMMATION

The immunosuppressive effects of the compositions of the invention against rheumatoid arthritis is determined in an experimental animal model system. The experimental model system is adjuvant induced arthritis in rats, and the protocol is described by J. Holoshitz, et at., 1983, Science, 219:56, or by B. Waksman et al., 1963, Int. Arch. Allergy Appl. Immunol., 23:129. Induction of the disease can be caused by a single injection, generally intradermally, of a suspension of killed Mycobacterium tuberculosis in complete Freund's adjuvant (CFA). The route of injection can vary, but rats may be injected at the base of the tail with an adjuvant mixture. The polypeptide is administered in phosphate buffered solution (PBS) at a dose of about 1-5 mg/kg. The control consists of administering PBS only.

The procedure for testing the effects of the test compound would consist of intradermally injecting killed Mycobacterium tuberculosis in CFA followed by immediately administering the test compound and subsequent treatment every other day until day 24. At 14, 15, 18, 20, 22, and 24 days after injection of Mycobacterium CFA, an overall arthritis score may be obtained as described by J. Holoskitz above. An analysis of the data would reveal that the test compound would have a dramatic affect on the swelling of the joints as measured by a decrease of the arthritis score.

4.11 THERAPEUTIC METHODS

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The compositions (including polypeptide fragments, analogs, variants and antibodies or other binding partners or modulators including antisense polynucleotides) of the invention have numerous applications in a variety of therapeutic methods. Examples of therapeutic applications include, but are not limited to, those exemplified herein.

4.11.1 EXAMPLE

One embodiment of the invention is the administration of an effective amount of the polypeptides or other composition of the invention to individuals affected by a disease or disorder that can be modulated by regulating the peptides of the invention. While the mode of administration is not particularly important, parenteral administration is preferred. An

exemplary mode of administration is to deliver an intravenous bolus. The dosage of the polypeptides or other composition of the invention will normally be determined by the prescribing physician. It is to be expected that the dosage will vary according to the age, weight, condition and response of the individual patient. Typically, the amount of polypeptide administered per dose will be in the range of about $0.01\mu g/kg$ to 100 mg/kg of body weight, with the preferred dose being about $0.1\mu g/kg$ to 10 mg/kg of patient body weight. For parenteral administration, polypeptides of the invention will be formulated in an injectable form combined with a pharmaceutically acceptable parenteral vehicle. Such vehicles are well known in the art and examples include water, saline, Ringer's solution, dextrose solution, and solutions consisting of small amounts of the human serum albumin. The vehicle may contain minor amounts of additives that maintain the isotonicity and stability of the polypeptide or other active ingredient. The preparation of such solutions is within the skill of the art.

15 4.12 PHARMACEUTICAL FORMULATIONS AND ROUTES OF ADMINISTRATION

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A protein or other composition of the present invention (from whatever source derived, including without limitation from recombinant and non-recombinant sources and including antibodies and other binding partners of the polypeptides of the invention) may be administered to a patient in need, by itself, or in pharmaceutical compositions where it is mixed with suitable carriers or excipient(s) at doses to treat or ameliorate a variety of disorders. Such a composition may optionally contain (in addition to protein or other active ingredient and a carrier) diluents, fillers, salts, buffers, stabilizers, solubilizers, and other materials well known in the art. The term "pharmaceutically acceptable" means a non-toxic material that does not interfere with the effectiveness of the biological activity of the active ingredient(s). The characteristics of the carrier will depend on the route of administration. The pharmaceutical composition of the invention may also contain cytokines, lymphokines, or other hematopoietic factors such as M-CSF, GM-CSF, TNF, IL-1, IL-2, IL-3, IL-4, IL-5, IL-6. IL-7. IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, IL-14, IL-15, IFN, TNF0, TNF1, TNF2, G-CSF, Meg-CSF, thrombopoietin, stem cell factor, and erythropoietin. In further compositions, proteins of the invention may be combined with other agents beneficial to the treatment of the disease or disorder in question. These agents include various growth factors such as epidermal growth factor (EGF), platelet-derived growth factor (PDGF), transforming

growth factors (TGF- α and TGF- β), insulin-like growth factor (IGF), as well as cytokines described herein.

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The pharmaceutical composition may further contain other agents which either enhance the activity of the protein or other active ingredient or complement its activity or use in treatment. Such additional factors and/or agents may be included in the pharmaceutical composition to produce a synergistic effect with protein or other active ingredient of the invention, or to minimize side effects. Conversely, protein or other active ingredient of the present invention may be included in formulations of the particular clotting factor, cytokine, lymphokine, other hematopoietic factor, thrombolytic or anti-thrombotic factor, or anti-inflammatory agent to minimize side effects of the clotting factor, cytokine, lymphokine, other hematopoietic factor, thrombolytic or anti-thrombotic factor, or anti-inflammatory agent (such as IL-1Ra, IL-1 Hy1, IL-1 Hy2, anti-TNF, corticosteroids, immunosuppressive agents). A protein of the present invention may be active in multimers (e.g., heterodimers or homodimers) or complexes with itself or other proteins. As a result, pharmaceutical compositions of the invention may comprise a protein of the invention in such multimeric or complexed form.

As an alternative to being included in a pharmaceutical composition of the invention including a first protein, a second protein or a therapeutic agent may be concurrently administered with the first protein (e.g., at the same time, or at differing times provided that therapeutic concentrations of the combination of agents is achieved at the treatment site). Techniques for formulation and administration of the compounds of the instant application may be found in "Remington's Pharmaceutical Sciences," Mack Publishing Co., Easton, PA, latest edition. A therapeutically effective dose further refers to that amount of the compound sufficient to result in amelioration of symptoms, e.g., treatment, healing, prevention or amelioration of the relevant medical condition, or an increase in rate of treatment, healing, prevention or amelioration of such conditions. When applied to an individual active ingredient, administered alone, a therapeutically effective dose refers to that ingredient alone. When applied to a combination, a therapeutically effective dose refers to combined amounts of the active ingredients that result in the therapeutic effect, whether administered in combination, serially or simultaneously.

In practicing the method of treatment or use of the present invention, a therapeutically effective amount of protein or other active ingredient of the present invention is administered to a mammal having a condition to be treated. Protein or other active ingredient of the

present invention may be administered in accordance with the method of the invention either alone or in combination with other therapies such as treatments employing cytokines, lymphokines or other hematopoietic factors. When co- administered with one or more cytokines, lymphokines or other hematopoietic factors, protein or other active ingredient of the present invention may be administered either simultaneously with the cytokine(s), lymphokine(s), other hematopoietic factor(s), thrombolytic or anti-thrombotic factors, or sequentially. If administered sequentially, the attending physician will decide on the appropriate sequence of administering protein or other active ingredient of the present invention in combination with cytokine(s), lymphokine(s), other hematopoietic factor(s), thrombolytic or anti-thrombotic factors.

4.12.1 ROUTES OF ADMINISTRATION

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Suitable routes of administration may, for example, include oral, rectal, transmucosal, or intestinal administration; parenteral delivery, including intramuscular, subcutaneous, intramedullary injections, as well as intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal, or intraocular injections. Administration of protein or other active ingredient of the present invention used in the pharmaceutical composition or to practice the method of the present invention can be carried out in a variety of conventional ways, such as oral ingestion, inhalation, topical application or cutaneous, subcutaneous, intraperitoneal, parenteral or intravenous injection. Intravenous administration to the patient is preferred.

Alternately, one may administer the compound in a local rather than systemic manner, for example, via injection of the compound directly into a arthritic joints or in fibrotic tissue, often in a depot or sustained release formulation. In order to prevent the scarring process frequently occurring as complication of glaucoma surgery, the compounds may be administered topically, for example, as eye drops. Furthermore, one may administer the drug in a targeted drug delivery system, for example, in a liposome coated with a specific antibody, targeting, for example, arthritic or fibrotic tissue. The liposomes will be targeted to and taken up selectively by the afflicted tissue.

The polypeptides of the invention are administered by any route that delivers an effective dosage to the desired site of action. The determination of a suitable route of administration and an effective dosage for a particular indication is within the level of skill in the art. Preferably for wound treatment, one administers the therapeutic compound directly to the site. Suitable dosage ranges for the polypeptides of the invention can be extrapolated

from these dosages or from similar studies in appropriate animal models. Dosages can then be adjusted as necessary by the clinician to provide maximal therapeutic benefit.

4.12.2 COMPOSITIONS/FORMULATIONS

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Pharmaceutical compositions for use in accordance with the present invention thus may be formulated in a conventional manner using one or more physiologically acceptable carriers comprising excipients and auxiliaries which facilitate processing of the active compounds into preparations which can be used pharmaceutically. These pharmaceutical compositions may be manufactured in a manner that is itself known, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilizing processes. Proper formulation is dependent upon the route of administration chosen. When a therapeutically effective amount of protein or other active ingredient of the present invention is administered orally, protein or other active ingredient of the present invention will be in the form of a tablet, capsule, powder, solution or elixir. When administered in tablet form, the pharmaceutical composition of the invention may additionally contain a solid carrier such as a gelatin or an adjuvant. The tablet, capsule, and powder contain from about 5 to 95% protein or other active ingredient of the present invention, and preferably from about 25 to 90% protein or other active ingredient of the present invention. When administered in liquid form, a liquid carrier such as water, petroleum, oils of animal or plant origin such as peanut oil, mineral oil, soybean oil, or sesame oil, or synthetic oils may be added. The liquid form of the pharmaceutical composition may further contain physiological saline solution, dextrose or other saccharide solution, or glycols such as ethylene glycol, propylene glycol or polyethylene glycol. When administered in liquid form, the pharmaceutical composition contains from about 0.5 to 90% by weight of protein or other active ingredient of the present invention, and preferably from about 1 to 50% protein or other active ingredient of the present invention.

When a therapeutically effective amount of protein or other active ingredient of the present invention is administered by intravenous, cutaneous or subcutaneous injection, protein or other active ingredient of the present invention will be in the form of a pyrogen-free, parenterally acceptable aqueous solution. The preparation of such parenterally acceptable protein or other active ingredient solutions, having due regard to pH, isotonicity, stability, and the like, is within the skill in the art. A preferred pharmaceutical composition for intravenous, cutaneous, or subcutaneous injection should contain, in addition to protein or

other active ingredient of the present invention, an isotonic vehicle such as Sodium Chloride Injection, Ringer's Injection, Dextrose Injection, Dextrose and Sodium Chloride Injection, Lactated Ringer's Injection, or other vehicle as known in the art. The pharmaceutical composition of the present invention may also contain stabilizers, preservatives, buffers, antioxidants, or other additives known to those of skill in the art. For injection, the agents of the invention may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hanks's solution, Ringer's solution, or physiological saline buffer. For transmucosal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

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For oral administration, the compounds can be formulated readily by combining the active compounds with pharmaceutically acceptable carriers well known in the art. Such carriers enable the compounds of the invention to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions and the like, for oral ingestion by a patient to be treated. Pharmaceutical preparations for oral use can be obtained from a solid excipient, optionally grinding a resulting mixture, and processing the mixture of granules, after adding suitable auxiliaries, if desired, to obtain tablets or dragee cores. Suitable excipients are, in particular, fillers such as sugars, including lactose, sucrose, mannitol, or sorbitol; cellulose preparations such as, for example, maize starch, wheat starch, rice starch, potato starch, gelatin, gum tragacanth, methyl cellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and/or polyvinylpyrrolidone (PVP). If desired, disintegrating agents may be added, such as the cross-linked polyvinyl pyrrolidone, agar, or alginic acid or a salt thereof such as sodium alginate. Dragee cores are provided with suitable coatings. For this purpose, concentrated sugar solutions may be used, which may optionally contain gum arabic, talc, polyvinyl pyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for identification or to characterize different combinations of active compound doses.

Pharmaceutical preparations which can be used orally include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a plasticizer, such as glycerol or sorbitol. The push-fit capsules can contain the active ingredients in admixture with filler such as lactose, binders such as starches, and/or lubricants such as talc or magnesium stearate and, optionally, stabilizers. In soft capsules, the active compounds may be dissolved or suspended in suitable liquids, such as fatty oils, liquid paraffin, or liquid polyethylene

glycols. In addition, stabilizers may be added. All formulations for oral administration should be in dosages suitable for such administration. For buccal administration, the compositions may take the form of tablets or lozenges formulated in conventional manner.

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For administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurized packs or a nebuliser, with the use of a suitable propellant, e.g., dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, e.g., gelatin for use in an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch. The compounds may be formulated for parenteral administration by injection, e.g., by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, e.g., in ampules or in multi-dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents.

Pharmaceutical formulations for parenteral administration include aqueous solutions of the active compounds in water-soluble form. Additionally, suspensions of the active compounds may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or synthetic fatty acid esters, such as ethyl oleate or triglycerides, or liposomes. Aqueous injection suspensions may contain substances which increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Optionally, the suspension may also contain suitable stabilizers or agents which increase the solubility of the compounds to allow for the preparation of highly concentrated solutions. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, e.g., sterile pyrogen-free water, before use.

The compounds may also be formulated in rectal compositions such as suppositories or retention enemas, e.g., containing conventional suppository bases such as cocoa butter or other glycerides. In addition to the formulations described previously, the compounds may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable

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polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

A pharmaceutical carrier for the hydrophobic compounds of the invention is a cosolvent system comprising benzyl alcohol, a nonpolar surfactant, a water-miscible organic polymer, and an aqueous phase. The co-solvent system may be the VPD co-solvent system. VPD is a solution of 3% w/v benzyl alcohol, 8% w/v of the nonpolar surfactant polysorbate 80, and 65% w/v polyethylene glycol 300, made up to volume in absolute ethanol. The VPD co-solvent system (VPD:5W) consists of VPD diluted 1:1 with a 5% dextrose in water solution. This co-solvent system dissolves hydrophobic compounds well, and itself produces low toxicity upon systemic administration. Naturally, the proportions of a co-solvent system may be varied considerably without destroying its solubility and toxicity characteristics. Furthermore, the identity of the co-solvent components may be varied; for example, other low-toxicity nonpolar surfactants may be used instead of polysorbate 80; the fraction size of polyethylene glycol may be varied; other biocompatible polymers may replace polyethylene glycol, e.g. polyvinyl pyrrolidone; and other sugars or polysaccharides may substitute for dextrose. Alternatively, other delivery systems for hydrophobic pharmaceutical compounds may be employed. Liposomes and emulsions are well known examples of delivery vehicles or carriers for hydrophobic drugs. Certain organic solvents such as dimethylsulfoxide also may be employed, although usually at the cost of greater toxicity. Additionally, the compounds may be delivered using a sustained-release system, such as semipermeable matrices of solid hydrophobic polymers containing the therapeutic agent. Various types of sustained-release materials have been established and are well known by those skilled in the art. Sustained-release capsules may, depending on their chemical nature, release the compounds for a few weeks up to over 100 days. Depending on the chemical nature and the biological stability of the therapeutic reagent, additional strategies for protein or other active ingredient stabilization may be employed.

The pharmaceutical compositions also may comprise suitable solid or gel phase carriers or excipients. Examples of such carriers or excipients include but are not limited to calcium carbonate, calcium phosphate, various sugars, starches, cellulose derivatives, gelatin, and polymers such as polyethylene glycols. Many of the active ingredients of the invention may be provided as salts with pharmaceutically compatible counter ions. Such pharmaceutically acceptable base addition salts are those salts which retain the biological effectiveness and properties of the free acids and which are obtained by reaction with

inorganic or organic bases such as sodium hydroxide, magnesium hydroxide, ammonia, trialkylamine, dialkylamine, monoalkylamine, dibasic amino acids, sodium acetate, potassium benzoate, triethanol amine and the like.

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The pharmaceutical composition of the invention may be in the form of a complex of the protein(s) or other active ingredient(s) of present invention along with protein or peptide antigens. The protein and/or peptide antigen will deliver a stimulatory signal to both B and T lymphocytes. B lymphocytes will respond to antigen through their surface immunoglobulin receptor. T lymphocytes will respond to antigen through the T cell receptor (TCR) following presentation of the antigen by MHC proteins. MHC and structurally related proteins including those encoded by class I and class II MHC genes on host cells will serve to present the peptide antigen(s) to T lymphocytes. The antigen components could also be supplied as purified MHC-peptide complexes alone or with co-stimulatory molecules that can directly signal T cells. Alternatively antibodies able to bind surface immunoglobulin and other molecules on B cells as well as antibodies able to bind the TCR and other molecules on T cells can be combined with the pharmaceutical composition of the invention.

The pharmaceutical composition of the invention may be in the form of a liposome in which protein of the present invention is combined, in addition to other pharmaceutically acceptable carriers, with amphipathic agents such as lipids which exist in aggregated form as micelles, insoluble monolayers, liquid crystals, or lamellar layers in aqueous solution. Suitable lipids for liposomal formulation include, without limitation, monoglycerides, diglycerides, sulfatides, lysolecithins, phospholipids, saponin, bile acids, and the like. Preparation of such liposomal formulations is within the level of skill in the art, as disclosed, for example, in U.S. Patent Nos. 4,235,871; 4,501,728; 4,837,028; and 4,737,323, all of which are incorporated herein by reference.

The amount of protein or other active ingredient of the present invention in the pharmaceutical composition of the present invention will depend upon the nature and severity of the condition being treated, and on the nature of prior treatments which the patient has undergone. Ultimately, the attending physician will decide the amount of protein or other active ingredient of the present invention with which to treat each individual patient. Initially, the attending physician will administer low doses of protein or other active ingredient of the present invention and observe the patient's response. Larger doses of protein or other active ingredient of the present invention may be administered until the optimal therapeutic effect is obtained for the patient, and at that point the dosage is not

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increased further. It is contemplated that the various pharmaceutical compositions used to practice the method of the present invention should contain about 0.01 µg to about 100 mg (preferably about 0.1 µg to about 10 mg, more preferably about 0.1 µg to about 1 mg) of protein or other active ingredient of the present invention per kg body weight. For compositions of the present invention which are useful for bone, cartilage, tendon or ligament regeneration, the therapeutic method includes administering the composition topically, systematically, or locally as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition may desirably be encapsulated or injected in a viscous form for delivery to the site of bone, cartilage or tissue damage. Topical administration may be suitable for wound healing and tissue repair. Therapeutically useful agents other than a protein or other active ingredient of the invention which may also optionally be included in the composition as described above, may alternatively or additionally, be administered simultaneously or sequentially with the composition in the methods of the invention. Preferably for bone and/or cartilage formation, the composition would include a matrix capable of delivering the protein-containing or other active ingredient-containing composition to the site of bone and/or cartilage damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. Such matrices may be formed of materials presently in use for other implanted medical applications.

The choice of matrix material is based on biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. The particular application of the compositions will define the appropriate formulation. Potential matrices for the compositions may be biodegradable and chemically defined calcium sulfate, tricalcium phosphate, hydroxyapatite, polylactic acid, polyglycolic acid and polyanhydrides. Other potential materials are biodegradable and biologically well-defined, such as bone or dermal collagen. Further matrices are comprised of pure proteins or extracellular matrix components. Other potential matrices are nonbiodegradable and chemically defined, such as sintered hydroxyapatite, bioglass, aluminates, or other ceramics. Matrices may be comprised of combinations of any of the above mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalcium phosphate. The bioceramics may be altered in composition, such as in calcium-aluminate-phosphate and processing to alter pore size, particle size, particle shape, and biodegradability. Presently preferred is a 50:50 (mole

weight) copolymer of lactic acid and glycolic acid in the form of porous particles having diameters ranging from 150 to 800 microns. In some applications, it will be useful to utilize a sequestering agent, such as carboxymethyl cellulose or autologous blood clot, to prevent the protein compositions from disassociating from the matrix.

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A preferred family of sequestering agents is cellulosic materials such as alkylcelluloses (including hydroxyalkylcelluloses), including methylcellulose, ethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, hydroxypropyl-methylcellulose, and carboxymethylcellulose, the most preferred being cationic salts of carboxymethylcellulose (CMC). Other preferred sequestering agents include hyaluronic acid, sodium alginate, poly(ethylene glycol), polyoxyethylene oxide, carboxyvinyl polymer and poly(vinyl alcohol). The amount of sequestering agent useful herein is 0.5-20 wt %, preferably 1-10 wt % based on total formulation weight, which represents the amount necessary to prevent desorption of the protein from the polymer matrix and to provide appropriate handling of the composition, yet not so much that the progenitor cells are prevented from infiltrating the matrix, thereby providing the protein the opportunity to assist the osteogenic activity of the progenitor cells. In further compositions, proteins or other active ingredients of the invention may be combined with other agents beneficial to the treatment of the bone and/or cartilage defect, wound, or tissue in question. These agents include various growth factors such as epidermal growth factor (EGF), platelet derived growth factor (PDGF), transforming growth factors (TGF- α and TGF- β), and insulin-like growth factor (IGF).

The therapeutic compositions are also presently valuable for veterinary applications. Particularly domestic animals and thoroughbred horses, in addition to humans, are desired patients for such treatment with proteins or other active ingredients of the present invention. The dosage regimen of a protein-containing pharmaceutical composition to be used in tissue regeneration will be determined by the attending physician considering various factors which modify the action of the proteins, *e.g.*, amount of tissue weight desired to be formed, the site of damage, the condition of the damaged tissue, the size of a wound, type of damaged tissue (*e.g.*, bone), the patient's age, sex, and diet, the severity of any infection, time of administration and other clinical factors. The dosage may vary with the type of matrix used in the reconstitution and with inclusion of other proteins in the pharmaceutical composition. For example, the addition of other known growth factors, such as IGF I (insulin like growth factor I), to the final composition, may also effect the dosage. Progress can be monitored by

periodic assessment of tissue/bone growth and/or repair, for example, X-rays, histomorphometric determinations and tetracycline labeling.

Polynucleotides of the present invention can also be used for gene therapy. Such polynucleotides can be introduced either in vivo or ex vivo into cells for expression in a mammalian subject. Polynucleotides of the invention may also be administered by other known methods for introduction of nucleic acid into a cell or organism (including, without limitation, in the form of viral vectors or naked DNA). Cells may also be cultured ex vivo in the presence of proteins of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes.

4.12.3 EFFECTIVE DOSAGE

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Pharmaceutical compositions suitable for use in the present invention include compositions wherein the active ingredients are contained in an effective amount to achieve its intended purpose. More specifically, a therapeutically effective amount means an amount effective to prevent development of or to alleviate the existing symptoms of the subject being treated. Determination of the effective amount is well within the capability of those skilled in the art, especially in light of the detailed disclosure provided herein. For any compound used in the method of the invention, the therapeutically effective dose can be estimated initially from appropriate in vitro assays. For example, a dose can be formulated in animal models to achieve a circulating concentration range that can be used to more accurately determine useful doses in humans. For example, a dose can be formulated in animal models to achieve a circulating concentration range that includes the IC₅₀ as determined in cell culture (i.e., the concentration of the test compound which achieves a half-maximal inhibition of the protein's biological activity). Such information can be used to more accurately determine useful doses in humans.

A therapeutically effective dose refers to that amount of the compound that results in amelioration of symptoms or a prolongation of survival in a patient. Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LD₅₀ (the dose lethal to 50% of the population) and the ED₅₀ (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio between LD₅₀ and ED₅₀. Compounds which exhibit high therapeutic

indices are preferred. The data obtained from these cell culture assays and animal studies can be used in formulating a range of dosage for use in human. The dosage of such compounds lies preferably within a range of circulating concentrations that include the ED₅₀ with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. The exact formulation, route of administration and dosage can be chosen by the individual physician in view of the patient's condition. See, e.g., Fingl et al., 1975, in "The Pharmacological Basis of Therapeutics", Ch. 1 p.1. Dosage amount and interval may be adjusted individually to provide plasma levels of the active moiety which are sufficient to maintain the desired effects, or minimal effective concentration (MEC). The MEC will vary for each compound but can be estimated from in vitro data. Dosages necessary to achieve the MEC will depend on individual characteristics and route of administration. However, HPLC assays or bioassays can be used to determine plasma concentrations.

Dosage intervals can also be determined using MEC value. Compounds should be administered using a regimen which maintains plasma levels above the MEC for 10-90% of the time, preferably between 30-90% and most preferably between 50-90%. In cases of local administration or selective uptake, the effective local concentration of the drug may not be related to plasma concentration.

An exemplary dosage regimen for polypeptides or other compositions of the invention will be in the range of about $0.01~\mu g/kg$ to 100~mg/kg of body weight daily, with the preferred dose being about $0.1~\mu g/kg$ to 25~mg/kg of patient body weight daily, varying in adults and children. Dosing may be once daily, or equivalent doses may be delivered at longer or shorter intervals.

The amount of composition administered will, of course, be dependent on the subject being treated, on the subject's age and weight, the severity of the affliction, the manner of administration and the judgment of the prescribing physician.

4.12.4 PACKAGING

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The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient. The pack may, for example, comprise metal or plastic foil, such as a blister pack. The pack or dispenser device may be accompanied by instructions for administration. Compositions comprising a compound of the invention formulated in a compatible pharmaceutical carrier may also be

prepared, placed in an appropriate container, and labeled for treatment of an indicated condition.

4.13 ANTIBODIES

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Also included in the invention are antibodies to proteins, or fragments of proteins of the invention. The term "antibody" as used herein refers to immunoglobulin molecules and immunologically active portions of immunoglobulin (Ig) molecules, i.e., molecules that contain an antigen binding site that specifically binds (immunoreacts with) an antigen. Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain, F_{ab} , and $F_{(ab')2}$ fragments, and an F_{ab} expression library. In general, an antibody molecule obtained from humans relates to any of the classes IgG, IgM, IgA, IgE and IgD, which differ from one another by the nature of the heavy chain present in the molecule. Certain classes have subclasses as well, such as IgG_1 , IgG_2 , and others. Furthermore, in humans, the light chain may be a kappa chain or a lambda chain. Reference herein to antibodies includes a reference to all such classes, subclasses and types of human antibody species.

An isolated related protein of the invention may be intended to serve as an antigen, or a portion or fragment thereof, and additionally can be used as an immunogen to generate antibodies that immunospecifically bind the antigen, using standard techniques for polyclonal and monoclonal antibody preparation. The full-length protein can be used or, alternatively, the invention provides antigenic peptide fragments of the antigen for use as immunogens. An antigenic peptide fragment comprises at least 6 amino acid residues of the amino acid sequence of the full length protein, such as the amino acid sequences shown in SEQ ID NO: 342-682, and encompasses an epitope thereof such that an antibody raised against the peptide forms a specific immune complex with the full length protein or with any fragment that contains the epitope. Preferably, the antigenic peptide comprises at least 10 amino acid residues, or at least 15 amino acid residues, or at least 20 amino acid residues, or at least 30 amino acid residues. Preferred epitopes encompassed by the antigenic peptide are regions of the protein that are located on its surface; commonly these are hydrophilic regions.

In certain embodiments of the invention, at least one epitope encompassed by the antigenic peptide is a region of -related protein that is located on the surface of the protein, e.g., a hydrophilic region. A hydrophobicity analysis of the human related protein sequence will indicate which regions of a related protein are particularly hydrophilic and, therefore, are likely to encode surface residues useful for targeting antibody production. As a means for

targeting antibody production, hydropathy plots showing regions of hydrophilicity and hydrophobicity may be generated by any method well known in the art, including, for example, the Kyte Doolittle or the Hopp Woods methods, either with or without Fourier transformation. See, *e.g.*, Hopp and Woods, 1981, *Proc. Nat. Acad. Sci. USA* 78: 3824-3828; Kyte and Doolittle 1982, *J. Mol. Biol.* 157: 105-142, each of which is incorporated herein by reference in its entirety. Antibodies that are specific for one or more domains within an antigenic protein, or derivatives, fragments, analogs or homologs thereof, are also provided herein.

A protein of the invention, or a derivative, fragment, analog, homolog or ortholog thereof, may be utilized as an immunogen in the generation of antibodies that immunospecifically bind these protein components.

Various procedures known within the art may be used for the production of polyclonal or monoclonal antibodies directed against a protein of the invention, or against derivatives, fragments, analogs homologs or orthologs thereof (see, for example, Antibodies: A Laboratory Manual, Harlow E, and Lane D, 1988, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, incorporated herein by reference). Some of these antibodies are discussed below.

4.13.1 POLYCLONAL ANTIBODIES

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For the production of polyclonal antibodies, various suitable host animals (e.g., rabbit, goat, mouse or other mammal) may be immunized by one or more injections with the native protein, a synthetic variant thereof, or a derivative of the foregoing. An appropriate immunogenic preparation can contain, for example, the naturally occurring immunogenic protein, a chemically synthesized polypeptide representing the immunogenic protein, or a recombinantly expressed immunogenic protein. Furthermore, the protein may be conjugated to a second protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. The preparation can further include an adjuvant. Various adjuvants used to increase the immunological response include, but are not limited to, Freund's (complete and incomplete), mineral gels (e.g., aluminum hydroxide), surface active substances (e.g., lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, dinitrophenol, etc.), adjuvants usable in humans such as Bacille Calmette-Guerin and Corynebacterium parvum, or similar immunostimulatory agents.

Additional examples of adjuvants which can be employed include MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate).

The polyclonal antibody molecules directed against the immunogenic protein can be isolated from the mammal (e.g., from the blood) and further purified by well known techniques, such as affinity chromatography using protein A or protein G, which provide primarily the IgG fraction of immune serum. Subsequently, or alternatively, the specific antigen which is the target of the immunoglobulin sought, or an epitope thereof, may be immobilized on a column to purify the immune specific antibody by immunoaffinity chromatography. Purification of immunoglobulins is discussed, for example, by D. Wilkinson (The Scientist, published by The Scientist, Inc., Philadelphia PA, Vol. 14, No. 8 (April 17, 2000), pp. 25-28).

4.13.2 MONOCLONAL ANTIBODIES

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The term "monoclonal antibody" (MAb) or "monoclonal antibody composition", as used herein, refers to a population of antibody molecules that contain only one molecular species of antibody molecule consisting of a unique light chain gene product and a unique heavy chain gene product. In particular, the complementarity determining regions (CDRs) of the monoclonal antibody are identical in all the molecules of the population. MAbs thus contain an antigen binding site capable of immunoreacting with a particular epitope of the antigen characterized by a unique binding affinity for it.

Monoclonal antibodies can be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes can be immunized in vitro.

The immunizing agent will typically include the protein antigen, a fragment thereof or a fusion protein thereof. Generally, either peripheral blood lymphocytes are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103). Immortalized cell lines are usually transformed mammalian cells, particularly

myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells can be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

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Preferred immortalized cell lines are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. More preferred immortalized cell lines are murine myeloma lines, which can be obtained, for instance, from the Salk Institute Cell Distribution Center, San Diego, California and the American Type Culture Collection, Manassas, Virginia. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); Brodeur et al., Monoclonal Antibody Production Techniques and Applications, Marcel Dekker, Inc., New York, (1987) pp. 51-63).

The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against the antigen. Preferably, the binding specificity of monoclonal antibodies produced by the hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA). Such techniques and assays are known in the art. The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson and Pollard, Anal. Biochem., 107:220 (1980). Preferably, antibodies having a high degree of specificity and a high binding affinity for the target antigen are isolated.

After the desired hybridoma cells are identified, the clones can be subcloned by limiting dilution procedures and grown by standard methods. Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium and RPMI-1640 medium. Alternatively, the hybridoma cells can be grown in vivo as ascites in a mammal. The monoclonal antibodies secreted by the subclones can be isolated or purified from the culture medium or ascites fluid by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

The monoclonal antibodies can also be made by recombinant DNA methods, such as those described in U.S. Patent No. 4,816,567. DNA encoding the monoclonal antibodies of the invention can be readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA can be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also can be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences (U.S. Patent No. 4,816,567; Morrison, Nature 368, 812-13 (1994)) or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a nonimmunoglobulin polypeptide. Such a non-immunoglobulin polypeptide can be substituted for the constant domains of an antibody of the invention, or can be substituted for the variable domains of one antigen-combining site of an antibody of the invention to create a chimeric bivalent antibody.

4.13.3 HUMANIZED ANTIBODIES

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The antibodies directed against the protein antigens of the invention can further comprise humanized antibodies or human antibodies. These antibodies are suitable for administration to humans without engendering an immune response by the human against the administered immunoglobulin. Humanized forms of antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')₂ or other antigen-binding subsequences of antibodies) that are principally comprised of the sequence of a human immunoglobulin, and contain minimal sequence derived from a non-human immunoglobulin. Humanization can be performed following the method of Winter and co-workers (Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. (See also U.S. Patent No. 5,225,539.) In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies can also comprise residues which are found neither in the recipient antibody nor in the

imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin (Jones et al., 1986; Riechmann et al., 1988; and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)).

4.13.4 HUMAN ANTIBODIES

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Fully human antibodies relate to antibody molecules in which essentially the entire sequences of both the light chain and the heavy chain, including the CDRs, arise from human genes. Such antibodies are termed "human antibodies", or "fully human antibodies" herein. Human monoclonal antibodies can be prepared by the trioma technique; the human B-cell hybridoma technique (see Kozbor, et al., 1983 Immunol Today 4: 72) and the EBV hybridoma technique to produce human monoclonal antibodies (see Cole, et al., 1985 In: MONOCLONAL ANTIBODIES AND CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96). Human monoclonal antibodies may be utilized in the practice of the present invention and may be produced by using human hybridomas (see Cote, et al., 1983. Proc Natl Acad Sci USA 80: 2026-2030) or by transforming human B-cells with Epstein Barr Virus in vitro (see Cole, et al., 1985 In: MONOCLONAL ANTIBODIES AND CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96).

In addition, human antibodies can also be produced using additional techniques, including phage display libraries (Hoogenboom and Winter, J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)). Similarly, human antibodies can be made by introducing human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in Marks et al. (Bio/Technology 10, 779-783 (1992)); Lonberg et al. (Nature 368 856-859 (1994)); Morrison (Nature 368, 812-13 (1994)); Fishwild et al.(Nature Biotechnology 14, 845-51 (1996)); Neuberger (Nature

Biotechnology 14, 826 (1996)); and Lonberg and Huszar (Intern. Rev. Immunol. 13 65-93 (1995)).

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Human antibodies may additionally be produced using transgenic nonhuman animals which are modified so as to produce fully human antibodies rather than the animal's endogenous antibodies in response to challenge by an antigen. (See PCT publication WO94/02602). The endogenous genes encoding the heavy and light immunoglobulin chains in the nonhuman host have been incapacitated, and active loci encoding human heavy and light chain immunoglobulins are inserted into the host's genome. The human genes are incorporated, for example, using yeast artificial chromosomes containing the requisite human DNA segments. An animal which provides all the desired modifications is then obtained as progeny by crossbreeding intermediate transgenic animals containing fewer than the full complement of the modifications. The preferred embodiment of such a nonhuman animal is a mouse, and is termed the XenomouseTM as disclosed in PCT publications WO 96/33735 and WO 96/34096. This animal produces B cells which secrete fully human immunoglobulins. The antibodies can be obtained directly from the animal after immunization with an immunogen of interest, as, for example, a preparation of a polyclonal antibody, or alternatively from immortalized B cells derived from the animal, such as hybridomas producing monoclonal antibodies. Additionally, the genes encoding the immunoglobulins with human variable regions can be recovered and expressed to obtain the antibodies directly, or can be further modified to obtain analogs of antibodies such as, for example, single chain Fv molecules.

An example of a method of producing a nonhuman host, exemplified as a mouse, lacking expression of an endogenous immunoglobulin heavy chain is disclosed in U.S. Patent No. 5,939,598. It can be obtained by a method including deleting the J segment genes from at least one endogenous heavy chain locus in an embryonic stem cell to prevent rearrangement of the locus and to prevent formation of a transcript of a rearranged immunoglobulin heavy chain locus, the deletion being effected by a targeting vector containing a gene encoding a selectable marker; and producing from the embryonic stem cell a transgenic mouse whose somatic and germ cells contain the gene encoding the selectable marker.

A method for producing an antibody of interest, such as a human antibody, is disclosed in U.S. Patent No. 5,916,771. It includes introducing an expression vector that contains a nucleotide sequence encoding a heavy chain into one mammalian host cell in

culture, introducing an expression vector containing a nucleotide sequence encoding a light chain into another mammalian host cell, and fusing the two cells to form a hybrid cell. The hybrid cell expresses an antibody containing the heavy chain and the light chain.

In a further improvement on this procedure, a method for identifying a clinically relevant epitope on an immunogen, and a correlative method for selecting an antibody that binds immunospecifically to the relevant epitope with high affinity, are disclosed in PCT publication WO 99/53049.

4.13.5 Fab FRAGMENTS AND SINGLE CHAIN ANTIBODIES

According to the invention, techniques can be adapted for the production of single-chain antibodies specific to an antigenic protein of the invention (see e.g., U.S. Patent No. 4,946,778). In addition, methods can be adapted for the construction of F_{ab} expression libraries (see e.g., Huse, et al., 1989 Science 246: 1275-1281) to allow rapid and effective identification of monoclonal F_{ab} fragments with the desired specificity for a protein or derivatives, fragments, analogs or homologs thereof. Antibody fragments that contain the idiotypes to a protein antigen may be produced by techniques known in the art including, but not limited to: (i) an $F_{(ab)2}$ fragment produced by pepsin digestion of an antibody molecule; (ii) an F_{ab} fragment generated by reducing the disulfide bridges of an $F_{(ab)2}$ fragment; (iii) an F_{ab} fragment generated by the treatment of the antibody molecule with papain and a reducing agent and (iv) F_{v} fragments.

4.13.6 BISPECIFIC ANTIBODIES

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Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for an antigenic protein of the invention. The second binding target is any other antigen, and advantageously is a cell-surface protein or receptor or receptor subunit.

Methods for making bispecific antibodies are known in the art. Traditionally, the recombinant production of bispecific antibodies is based on the co-expression of two immunoglobulin heavy-chain/light-chain pairs, where the two heavy chains have different specificities (Milstein and Cuello, Nature, 305:537-539 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of ten different antibody molecules, of which only one has the

correct bispecific structure. The purification of the correct molecule is usually accomplished by affinity chromatography steps. Similar procedures are disclosed in WO 93/08829, published 13 May 1993, and in Traunecker *et al.*, 1991 *EMBO J.*, 10:3655-3659.

Antibody variable domains with the desired binding specificities (antibody-antigen combining sites) can be fused to immunoglobulin constant domain sequences. The fusion preferably is with an immunoglobulin heavy-chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1) containing the site necessary for light-chain binding present in at least one of the fusions. DNAs encoding the immunoglobulin heavy-chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are cotransfected into a suitable host organism. For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology, 121:210 (1986).

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According to another approach described in WO 96/27011, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the CH3 region of an antibody constant domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). This provides a mechanism for increasing the yield of the heterodimer over other unwanted end-products such as homodimers.

Bispecific antibodies can be prepared as full length antibodies or antibody fragments (e.g. F(ab')₂ bispecific antibodies). Techniques for generating bispecific antibodies from antibody fragments have been described in the literature. For example, bispecific antibodies can be prepared using chemical linkage. Brennan et al., Science 229:81 (1985) describe a procedure wherein intact antibodies are proteolytically cleaved to generate F(ab')₂ fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB

derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

Additionally, Fab' fragments can be directly recovered from E. coli and chemically coupled to form bispecific antibodies. Shalaby et al., J. Exp. Med. 175:217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab')₂ molecule. Each Fab' fragment was separately secreted from E. coli and subjected to directed chemical coupling in vitro to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the ErbB2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets.

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Various techniques for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelny et al., J. Immunol. 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., Proc. Natl. Acad. Sci. USA 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a heavy-chain variable domain (V_H) connected to a light-chain variable domain (V_L) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the V_H and V_L domains of one fragment are forced to pair with the complementary V_L and V_H domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See, Gruber et al., J. Immunol. 152:5368 (1994).

Antibodies with more than two valencies are contemplated. For example, trispecific antibodies can be prepared. Tutt et al., <u>J. Immunol.</u> 147:60 (1991). Exemplary bispecific antibodies can bind to two different epitopes, at least one of which originates in the protein antigen of the invention. Alternatively, an anti-antigenic arm of an immunoglobulin molecule can be combined with an arm which binds to a triggering molecule on a leukocyte such as a T-cell receptor molecule (e.g. CD2, CD3, CD28, or B7), or Fc receptors for IgG (FcγR), such as FcγRI (CD64), FcγRII (CD32) and FcγRIII (CD16) so as to focus cellular defense mechanisms to the cell expressing the particular antigen. Bispecific

antibodies can also be used to direct cytotoxic agents to cells which express a particular antigen. These antibodies possess an antigen-binding arm and an arm which binds a cytotoxic agent or a radionuclide chelator, such as EOTUBE, DPTA, DOTA, or TETA. Another bispecific antibody of interest binds the protein antigen described herein and further binds tissue factor (TF).

4.13.7 HETEROCONJUGATE ANTIBODIES

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Heteroconjugate antibodies are also within the scope of the present invention. Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (U.S. Patent No. 4,676,980), and for treatment of HIV infection (WO 91/00360; WO 92/200373; EP 03089). It is contemplated that the antibodies can be prepared in vitro using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins can be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminothiolate and methyl-4-mercaptobutyrimidate and those disclosed, for example, in U.S. Patent No. 4,676,980.

4.13.8 EFFECTOR FUNCTION ENGINEERING

It can be desirable to modify the antibody of the invention with respect to effector function, so as to enhance, e.g., the effectiveness of the antibody in treating cancer. For example, cysteine residue(s) can be introduced into the Fc region, thereby allowing interchain disulfide bond formation in this region. The homodimeric antibody thus generated can have improved internalization capability and/or increased complement-mediated cell killing and antibody-dependent cellular cytotoxicity (ADCC). See Caron et al., J. Exp Med., 176: 1191-1195 (1992) and Shopes, J. Immunol., 148: 2918-2922 (1992). Homodimeric antibodies with enhanced anti-tumor activity can also be prepared using heterobifunctional cross-linkers as described in Wolff et al. Cancer Research, 53: 2560-2565 (1993). Alternatively, an antibody can be engineered that has dual Fc regions and can thereby have enhanced complement lysis and ADCC capabilities. See Stevenson et al., Anti-Cancer Drug Design, 3: 219-230 (1989).

4.13.9 IMMUNOCONJUGATES

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The invention also pertains to immunoconjugates comprising an antibody conjugated to a cytotoxic agent such as a chemotherapeutic agent, toxin (e.g., an enzymatically active toxin of bacterial, fungal, plant, or animal origin, or fragments thereof), or a radioactive isotope (i.e., a radioconjugate).

Chemotherapeutic agents useful in the generation of such immunoconjugates have been described above. Enzymatically active toxins and fragments thereof that can be used include diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from Pseudomonas aeruginosa), ricin A chain, abrin A chain, modeccin A chain, alpha-sarcin, Aleurites fordii proteins, dianthin proteins, Phytolaca americana proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, crotin, sapaonaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin, and the tricothecenes. A variety of radionuclides are available for the production of radioconjugated antibodies. Examples include ²¹²Bi, ¹³¹I, ¹³¹In, ⁹⁰Y, and ¹⁸⁶Re.

Conjugates of the antibody and cytotoxic agent are made using a variety of bifunctional protein-coupling agents such as N-succinimidyl-3-(2-pyridyldithiol) propionate (SPDP), iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutareldehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexanediamine), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as tolyene 2,6-diisocyanate), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, a ricin immunotoxin can be prepared as described in Vitetta et al., Science, 238: 1098 (1987). Carbon-14-labeled 1-isothiocyanatobenzyl-3-methyldiethylene triaminepentaacetic acid (MX-DTPA) is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026.

In another embodiment, the antibody can be conjugated to a "receptor" (such as streptavidin) for utilization in tumor pretargeting wherein the antibody-receptor conjugate is administered to the patient, followed by removal of unbound conjugate from the circulation using a clearing agent and then administration of a "ligand" (e.g., avidin) that is in turn conjugated to a cytotoxic agent.

4.14 COMPUTER READABLE SEQUENCES

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In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any medium which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A skilled artisan can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising computer readable medium having recorded thereon a nucleotide sequence of the present invention. As used herein, "recorded" refers to a process for storing information on computer readable medium. A skilled artisan can readily adopt any of the presently known methods for recording information on computer readable medium to generate manufactures comprising the nucleotide sequence information of the present invention.

A variety of data storage structures are available to a skilled artisan for creating a computer readable medium having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the present invention on computer readable medium. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and Microsoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A skilled artisan can readily adapt any number of data processor structuring formats (e.g. text file or database) in order to obtain computer readable medium having recorded thereon the nucleotide sequence information of the present invention.

By providing any of the nucleotide sequences SEQ ID NO: 1-341 or a representative fragment thereof; or a nucleotide sequence at least 95% identical to any of the nucleotide sequences of SEQ ID NO: 1-341 in computer readable form, a skilled artisan can routinely access the sequence information for a variety of purposes. Computer software is publicly available which allows a skilled artisan to access sequence information provided in a computer readable medium. The examples which follow demonstrate how software which implements the BLAST (Altschul et al., J. Mol. Biol. 215:403-410 (1990)) and BLAZE

(Brutlag et al., Comp. Chem. 17:203-207 (1993)) search algorithms on a Sybase system is used to identify open reading frames (ORFs) within a nucleic acid sequence. Such ORFs may be protein encoding fragments and may be useful in producing commercially important proteins such as enzymes used in fermentation reactions and in the production of commercially useful metabolites.

As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based systems are suitable for use in the present invention. As stated above, the computer-based systems of the present invention comprise a data storage means having stored therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means. As used herein, "data storage means" refers to memory which can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention.

As used herein, "search means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage means. Search means are used to identify fragments or regions of a known sequence which match a particular target sequence or target motif. A variety of known algorithms are disclosed publicly and a variety of commercially available software for conducting search means are and can be used in the computer-based systems of the present invention. Examples of such software includes, but is not limited to, Smith-Waterman, MacPattern (EMBL), BLASTN and BLASTA (NPOLYPEPTIDEIA). A skilled artisan can readily recognize that any one of the available algorithms or implementing software packages for conducting homology searches can be adapted for use in the present computer-based systems. As used herein, a "target sequence" can be any nucleic acid or amino acid sequence of six or more nucleotides or two or more amino acids. A skilled artisan can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 300 amino acids,

more preferably from about 30 to 100 nucleotide residues. However, it is well recognized that searches for commercially important fragments, such as sequence fragments involved in gene expression and protein processing, may be of shorter length.

As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration which is formed upon the folding of the target motif. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzyme active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

4.15 TRIPLE HELIX FORMATION

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In addition, the fragments of the present invention, as broadly described, can be used to control gene expression through triple helix formation or antisense DNA or RNA, both of which methods are based on the binding of a polynucleotide sequence to DNA or RNA. Polynucleotides suitable for use in these methods are preferably 20 to 40 bases in length and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 15241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Olmno, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide.

4.16 DIAGNOSTIC ASSAYS AND KITS

The present invention further provides methods to identify the presence or expression of one of the ORFs of the present invention, or homolog thereof, in a test sample, using a nucleic acid probe or antibodies of the present invention, optionally conjugated or otherwise associated with a suitable label.

In general, methods for detecting a polynucleotide of the invention can comprise contacting a sample with a compound that binds to and forms a complex with the

polynucleotide for a period sufficient to form the complex, and detecting the complex, so that if a complex is detected, a polynucleotide of the invention is detected in the sample. Such methods can also comprise contacting a sample under stringent hybridization conditions with nucleic acid primers that anneal to a polynucleotide of the invention under such conditions, and amplifying annealed polynucleotides, so that if a polynucleotide is amplified, a polynucleotide of the invention is detected in the sample.

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In general, methods for detecting a polypeptide of the invention can comprise contacting a sample with a compound that binds to and forms a complex with the polypeptide for a period sufficient to form the complex, and detecting the complex, so that if a complex is detected, a polypeptide of the invention is detected in the sample.

In detail, such methods comprise incubating a test sample with one or more of the antibodies or one or more of the nucleic acid probes of the present invention and assaying for binding of the nucleic acid probes or antibodies to components within the test sample.

Conditions for incubating a nucleic acid probe or antibody with a test sample vary. Incubation conditions depend on the format employed in the assay, the detection methods employed, and the type and nature of the nucleic acid probe or antibody used in the assay. One skilled in the art will recognize that any one of the commonly available hybridization, amplification or immunological assay formats can readily be adapted to employ the nucleic acid probes or antibodies of the present invention. Examples of such assays can be found in Chard, T., An Introduction to Radioimmunoassay and Related Techniques, Elsevier Science Publishers, Amsterdam, The Netherlands (1986); Bullock, G.R. et al., Techniques in Immunocytochemistry, Academic Press, Orlando, FL Vol. 1 (1982), Vol. 2 (1983), Vol. 3 (1985); Tijssen, P., Practice and Theory of immunoassays: Laboratory Techniques in Biochemistry and Molecular Biology, Elsevier Science Publishers, Amsterdam, The Netherlands (1985). The test samples of the present invention include cells, protein or membrane extracts of cells, or biological fluids such as sputum, blood, serum, plasma, or urine. The test sample used in the above-described method will vary based on the assay format, nature of the detection method and the tissues, cells or extracts used as the sample to be assayed. Methods for preparing protein extracts or membrane extracts of cells are well known in the art and can be readily be adapted in order to obtain a sample which is compatible with the system utilized.

In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the assays of the present invention. Specifically, the

invention provides a compartment kit to receive, in close confinement, one or more containers which comprises: (a) a first container comprising one of the probes or antibodies of the present invention; and (b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of a bound probe or antibody.

In detail, a compartment kit includes any kit in which reagents are contained in separate containers. Such containers include small glass containers, plastic containers or strips of plastic or paper. Such containers allows one to efficiently transfer reagents from one compartment to another compartment such that the samples and reagents are not cross-contaminated, and the agents or solutions of each container can be added in a quantitative fashion from one compartment to another. Such containers will include a container which will accept the test sample, a container which contains the antibodies used in the assay, containers which contain wash reagents (such as phosphate buffered saline, Tris-buffers, etc.), and containers which contain the reagents used to detect the bound antibody or probe. Types of detection reagents include labeled nucleic acid probes, labeled secondary antibodies, or in the alternative, if the primary antibody is labeled, the enzymatic, or antibody binding reagents which are capable of reacting with the labeled antibody. One skilled in the art will readily recognize that the disclosed probes and antibodies of the present invention can be readily incorporated into one of the established kit formats which are well known in the art.

4.17 MEDICAL IMAGING

The novel polypeptides and binding partners of the invention are useful in medical imaging of sites expressing the molecules of the invention (e.g., where the polypeptide of the invention is involved in the immune response, for imaging sites of inflammation or infection). See, e.g., Kunkel et al., U.S. Pat. NO. 5,413,778. Such methods involve chemical attachment of a labeling or imaging agent, administration of the labeled polypeptide to a subject in a pharmaceutically acceptable carrier, and imaging the labeled polypeptide *in vivo* at the target site.

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4.18 SCREENING ASSAYS

Using the isolated proteins and polynucleotides of the invention, the present invention further provides methods of obtaining and identifying agents which bind to a polypeptide

encoded by an ORF corresponding to any of the nucleotide sequences set forth in SEQ ID NO: 1-341, or bind to a specific domain of the polypeptide encoded by the nucleic acid. In detail, said method comprises the steps of:

(a) contacting an agent with an isolated protein encoded by an ORF of the present invention, or nucleic acid of the invention; and

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(b) determining whether the agent binds to said protein or said nucleic acid.

In general, therefore, such methods for identifying compounds that bind to a polynucleotide of the invention can comprise contacting a compound with a polynucleotide of the invention for a time sufficient to form a polynucleotide/compound complex, and detecting the complex, so that if a polynucleotide/compound complex is detected, a compound that binds to a polynucleotide of the invention is identified.

Likewise, in general, therefore, such methods for identifying compounds that bind to a polypeptide of the invention can comprise contacting a compound with a polypeptide of the invention for a time sufficient to form a polypeptide/compound complex, and detecting the complex, so that if a polypeptide/compound complex is detected, a compound that binds to a polynucleotide of the invention is identified.

Methods for identifying compounds that bind to a polypeptide of the invention can also comprise contacting a compound with a polypeptide of the invention in a cell for a time sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a receptor gene sequence in the cell, and detecting the complex by detecting reporter gene sequence expression, so that if a polypeptide/compound complex is detected, a compound that binds a polypeptide of the invention is identified.

Compounds identified via such methods can include compounds which modulate the activity of a polypeptide of the invention (that is, increase or decrease its activity, relative to activity observed in the absence of the compound). Alternatively, compounds identified via such methods can include compounds which modulate the expression of a polynucleotide of the invention (that is, increase or decrease expression relative to expression levels observed in the absence of the compound). Compounds, such as compounds identified via the methods of the invention, can be tested using standard assays well known to those of skill in the art for their ability to modulate activity/expression.

The agents screened in the above assay can be, but are not limited to, peptides, carbohydrates, vitamin derivatives, or other pharmaceutical agents. The agents can be

selected and screened at random or rationally selected or designed using protein modeling techniques.

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For random screening, agents such as peptides, carbohydrates, pharmaceutical agents and the like are selected at random and are assayed for their ability to bind to the protein encoded by the ORF of the present invention. Alternatively, agents may be rationally selected or designed. As used herein, an agent is said to be "rationally selected or designed" when the agent is chosen based on the configuration of the particular protein. For example, one skilled in the art can readily adapt currently available procedures to generate peptides, pharmaceutical agents and the like, capable of binding to a specific peptide sequence, in order to generate rationally designed antipeptide peptides, for example see Hurby et al., Application of Synthetic Peptides: Antisense Peptides," In Synthetic Peptides, A User's Guide, W.H. Freeman, NY (1992), pp. 289-307, and Kaspczak et al., Biochemistry 28:9230-8 (1989), or pharmaceutical agents, or the like.

In addition to the foregoing, one class of agents of the present invention, as broadly described, can be used to control gene expression through binding to one of the ORFs or EMFs of the present invention. As described above, such agents can be randomly screened or rationally designed/selected. Targeting the ORF or EMF allows a skilled artisan to design sequence specific or element specific agents, modulating the expression of either a single ORF or multiple ORFs which rely on the same EMF for expression control. One class of DNA binding agents are agents which contain base residues which hybridize or form a triple helix formation by binding to DNA or RNA. Such agents can be based on the classic phosphodiester, ribonucleic acid backbone, or can be a variety of sulfhydryl or polymeric derivatives which have base attachment capacity.

Agents suitable for use in these methods preferably contain 20 to 40 bases and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Okano, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present

invention is necessary for the design of an antisense or triple helix oligonucleotide and other DNA binding agents.

Agents which bind to a protein encoded by one of the ORFs of the present invention can be used as a diagnostic agent. Agents which bind to a protein encoded by one of the ORFs of the present invention can be formulated using known techniques to generate a pharmaceutical composition.

4.19 USE OF NUCLEIC ACIDS AS PROBES

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Another aspect of the subject invention is to provide for polypeptide-specific nucleic acid hybridization probes capable of hybridizing with naturally occurring nucleotide sequences. The hybridization probes of the subject invention may be derived from any of the nucleotide sequences SEQ ID NO: 1-341. Because the corresponding gene is only expressed in a limited number of tissues, a hybridization probe derived from any of the nucleotide sequences SEQ ID NO: 1-341 can be used as an indicator of the presence of RNA of cell type of such a tissue in a sample.

Any suitable hybridization technique can be employed, such as, for example, in situ hybridization. PCR as described in US Patents Nos. 4,683,195 and 4,965,188 provides additional uses for oligonucleotides based upon the nucleotide sequences. Such probes used in PCR may be of recombinant origin, may be chemically synthesized, or a mixture of both. The probe will comprise a discrete nucleotide sequence for the detection of identical sequences or a degenerate pool of possible sequences for identification of closely related genomic sequences.

Other means for producing specific hybridization probes for nucleic acids include the cloning of nucleic acid sequences into vectors for the production of mRNA probes. Such vectors are known in the art and are commercially available and may be used to synthesize RNA probes *in vitro* by means of the addition of the appropriate RNA polymerase as T7 or SP6 RNA polymerase and the appropriate radioactively labeled nucleotides. The nucleotide sequences may be used to construct hybridization probes for mapping their respective genomic sequences. The nucleotide sequence provided herein may be mapped to a chromosome or specific regions of a chromosome using well known genetic and/or chromosomal mapping techniques. These techniques include in situ hybridization, linkage analysis against known chromosomal markers, hybridization screening with libraries or flow-sorted chromosomal preparations specific to known chromosomes, and the like. The

technique of fluorescent in situ hybridization of chromosome spreads has been described, among other places, in Verma et al (1988) Human Chromosomes: A Manual of Basic Techniques, Pergamon Press, New York NY.

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Fluorescent *in situ* hybridization of chromosomal preparations and other physical chromosome mapping techniques may be correlated with additional genetic map data. Examples of genetic map data can be found in the 1994 Genome Issue of Science (265:1981f). Correlation between the location of a nucleic acid on a physical chromosomal map and a specific disease (or predisposition to a specific disease) may help delimit the region of DNA associated with that genetic disease. The nucleotide sequences of the subject invention may be used to detect differences in gene sequences between normal, carrier or affected individuals.

4.20 PREPARATION OF SUPPORT BOUND OLIGONUCLEOTIDES

Oligonucleotides, i.e., small nucleic acid segments, may be readily prepared by, for example, directly synthesizing the oligonucleotide by chemical means, as is commonly practiced using an automated oligonucleotide synthesizer.

Support bound oligonucleotides may be prepared by any of the methods known to those of skill in the art using any suitable support such as glass, polystyrene or Teflon. One strategy is to precisely spot oligonucleotides synthesized by standard synthesizers. Immobilization can be achieved using passive adsorption (Inouye & Hondo, (1990) J. Clin. Microbiol. 28(6) 1469-72); using UV light (Nagata et al., 1985; Dahlen et al., 1987; Morrissey & Collins, (1989) Mol. Cell Probes 3(2) 189-207) or by covalent binding of base modified DNA (Keller et al., 1988; 1989); all references being specifically incorporated herein.

Another strategy that may be employed is the use of the strong biotin-streptavidin interaction as a linker. For example, Broude *et al.* (1994) Proc. Natl. Acad. Sci. USA 91(8) 3072-6, describe the use of biotinylated probes, although these are duplex probes, that are immobilized on streptavidin-coated magnetic beads. Streptavidin-coated beads may be purchased from Dynal, Oslo. Of course, this same linking chemistry is applicable to coating any surface with streptavidin. Biotinylated probes may be purchased from various sources, such as, e.g., Operon Technologies (Alameda, CA).

Nunc Laboratories (Naperville, IL) is also selling suitable material that could be used. Nunc Laboratories have developed a method by which DNA can be covalently bound to the microwell surface termed Covalink NH. CovaLink NH is a polystyrene surface grafted with

secondary amino groups (>NH) that serve as bridge-heads for further covalent coupling. CovaLink Modules may be purchased from Nunc Laboratories. DNA molecules may be bound to CovaLink exclusively at the 5'-end by a phosphoramidate bond, allowing immobilization of more than 1 pmol of DNA (Rasmussen *et al.*, (1991) Anal. Biochem. 198(1) 138-42).

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The use of CovaLink NH strips for covalent binding of DNA molecules at the 5'-end has been described (Rasmussen et al., (1991). In this technology, a phosphoramidate bond is employed (Chu et al., (1983) Nucleic Acids Res. 11(8) 6513-29). This is beneficial as immobilization using only a single covalent bond is preferred. The phosphoramidate bond joins the DNA to the CovaLink NH secondary amino groups that are positioned at the end of spacer arms covalently grafted onto the polystyrene surface through a 2 nm long spacer arm. To link an oligonucleotide to CovaLink NH via an phosphoramidate bond, the oligonucleotide terminus must have a 5'-end phosphate group. It is, perhaps, even possible for biotin to be covalently bound to CovaLink and then streptavidin used to bind the probes.

More specifically, the linkage method includes dissolving DNA in water (7.5 ng/μl) and denaturing for 10 min. at 95°C and cooling on ice for 10 min. Ice-cold 0.1 M 1-methylimidazole, pH 7.0 (1-MeIm₇), is then added to a final concentration of 10 mM 1-MeIm₇. The single-stranded DNA solution is then dispensed into CovaLink NH strips (75 μl/well) standing on ice.

Carbodiimide 0.2 M 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC), dissolved in 10 mM 1-MeIm₇, is made fresh and 25 µl added per well. The strips are incubated for 5 hours at 50°C. After incubation the strips are washed using, e.g., Nunc-Immuno Wash; first the wells are washed 3 times, then they are soaked with washing solution for 5 min., and finally they are washed 3 times (where in the washing solution is 0.4 N NaOH, 0.25% SDS heated to 50°C).

It is contemplated that a further suitable method for use with the present invention is that described in PCT Patent Application WO 90/03382 (Southern & Maskos), incorporated herein by reference. This method of preparing an oligonucleotide bound to a support involves attaching a nucleoside 3'-reagent through the phosphate group by a covalent phosphodiester link to aliphatic hydroxyl groups carried by the support. The oligonucleotide is then synthesized on the supported nucleoside and protecting groups removed from the synthetic oligonucleotide chain under standard conditions that do not cleave the oligonucleotide from the support. Suitable reagents include nucleoside phosphoramidite and nucleoside hydrogen phosphorate.

An on-chip strategy for the preparation of DNA probe for the preparation of DNA probe arrays may be employed. For example, addressable laser-activated photodeprotection may be

employed in the chemical synthesis of oligonucleotides directly on a glass surface, as described by Fodor *et al.* (1991) Science 251(4995) 767-73, incorporated herein by reference. Probes may also be immobilized on nylon supports as described by Van Ness *et al.* (1991) Nucleic Acids Res. 19(12) 3345-50; or linked to Teflon using the method of Duncan & Cavalier (1988) Anal. Biochem. 169(1) 104-8; all references being specifically incorporated herein.

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To link an oligonucleotide to a nylon support, as described by Van Ness *et al.* (1991), requires activation of the nylon surface via alkylation and selective activation of the 5'-amine of oligonucleotides with cyanuric chloride.

One particular way to prepare support bound oligonucleotides is to utilize the light-generated synthesis described by Pease *et al.*, (1994) PNAS USA 91(11) 5022-6, incorporated herein by reference). These authors used current photolithographic techniques to generate arrays of immobilized oligonucleotide probes (DNA chips). These methods, in which light is used to direct the synthesis of oligonucleotide probes in high-density, miniaturized arrays, utilize photolabile 5'-protected *N*-acyl-deoxynucleoside phosphoramidites, surface linker chemistry and versatile combinatorial synthesis strategies. A matrix of 256 spatially defined oligonucleotide probes may be generated in this manner.

4.21 PREPARATION OF NUCLEIC ACID FRAGMENTS

The nucleic acids may be obtained from any appropriate source, such as cDNAs, genomic DNA, chromosomal DNA, microdissected chromosome bands, cosmid or YAC inserts, and RNA, including mRNA without any amplification steps. For example, Sambrook *et al.* (1989) describes three protocols for the isolation of high molecular weight DNA from mammalian cells (p. 9.14-9.23).

DNA fragments may be prepared as clones in M13, plasmid or lambda vectors and/or prepared directly from genomic DNA or cDNA by PCR or other amplification methods. Samples may be prepared or dispensed in multiwell plates. About 100-1000 ng of DNA samples may be prepared in 2-500 ml of final volume.

The nucleic acids would then be fragmented by any of the methods known to those of skill in the art including, for example, using restriction enzymes as described at 9.24-9.28 of Sambrook *et al.* (1989), shearing by ultrasound and NaOH treatment.

Low pressure shearing is also appropriate, as described by Schriefer *et al.* (1990) Nucleic Acids Res. 18(24) 7455-6, incorporated herein by reference). In this method, DNA samples are passed through a small French pressure cell at a variety of low to intermediate pressures. A

lever device allows controlled application of low to intermediate pressures to the cell. The results of these studies indicate that low-pressure shearing is a useful alternative to sonic and enzymatic DNA fragmentation methods.

One particularly suitable way for fragmenting DNA is contemplated to be that using the two base recognition endonuclease, *CviJI*, described by Fitzgerald *et al.* (1992) Nucleic Acids Res. 20(14) 3753-62. These authors described an approach for the rapid fragmentation and fractionation of DNA into particular sizes that they contemplated to be suitable for shotgun cloning and sequencing.

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The restriction endonuclease CviII normally cleaves the recognition sequence PuGCPy between the G and C to leave blunt ends. Atypical reaction conditions, which alter the specificity of this enzyme ($CviII^{**}$), yield a quasi-random distribution of DNA fragments form the small molecule pUC19 (2688 base pairs). Fitzgerald $et\ al$. (1992) quantitatively evaluated the randomness of this fragmentation strategy, using a $CviII^{**}$ digest of pUC19 that was size fractionated by a rapid gel filtration method and directly ligated, without end repair, to a lac Z minus M13 cloning vector. Sequence analysis of 76 clones showed that $CviII^{**}$ restricts pyGCPy and PuGCPu, in addition to PuGCPy sites, and that new sequence data is accumulated at a rate consistent with random fragmentation.

As reported in the literature, advantages of this approach compared to sonication and agarose gel fractionation include: smaller amounts of DNA are required (0.2-0.5 µg instead of 2-5 µg); and fewer steps are involved (no preligation, end repair, chemical extraction, or agarose gel electrophoresis and elution are needed

Irrespective of the manner in which the nucleic acid fragments are obtained or prepared, it is important to denature the DNA to give single stranded pieces available for hybridization. This is achieved by incubating the DNA solution for 2-5 minutes at 80-90°C. The solution is then cooled quickly to 2°C to prevent renaturation of the DNA fragments before they are contacted with the chip. Phosphate groups must also be removed from genomic DNA by methods known in the art.

4.22 PREPARATION OF DNA ARRAYS

Arrays may be prepared by spotting DNA samples on a support such as a nylon membrane. Spotting may be performed by using arrays of metal pins (the positions of which correspond to an array of wells in a microtiter plate) to repeated by transfer of about 20 nl of a DNA solution to a nylon membrane. By offset printing, a density of dots higher than the density

of the wells is achieved. One to 25 dots may be accommodated in 1 mm², depending on the type of label used. By avoiding spotting in some preselected number of rows and columns, separate subsets (subarrays) may be formed. Samples in one subarray may be the same genomic segment of DNA (or the same gene) from different individuals, or may be different, overlapped genomic clones. Each of the subarrays may represent replica spotting of the same samples. In one example, a selected gene segment may be amplified from 64 patients. For each patient, the amplified gene segment may be in one 96-well plate (all 96 wells containing the same sample). A plate for each of the 64 patients is prepared. By using a 96-pin device, all samples may be spotted on one 8 x 12 cm membrane. Subarrays may contain 64 samples, one from each patient. Where the 96 subarrays are identical, the dot span may be 1 mm² and there may be a 1 mm space between subarrays.

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Another approach is to use membranes or plates (available from NUNC, Naperville, Illinois) which may be partitioned by physical spacers e.g. a plastic grid molded over the membrane, the grid being similar to the sort of membrane applied to the bottom of multiwell plates, or hydrophobic strips. A fixed physical spacer is not preferred for imaging by exposure to flat phosphor-storage screens or x-ray films.

The present invention is illustrated in the following examples. Upon consideration of the present disclosure, one of skill in the art will appreciate that many other embodiments and variations may be made in the scope of the present invention. Accordingly, it is intended that the broader aspects of the present invention not be limited to the disclosure of the following examples. The present invention is not to be limited in scope by the exemplified embodiments which are intended as illustrations of single aspects of the invention, and compositions and methods which are functionally equivalent are within the scope of the invention. Indeed, numerous modifications and variations in the practice of the invention are expected to occur to those skilled in the art upon consideration of the present preferred embodiments. Consequently, the only limitations which should be placed upon the scope of the invention are those which appear in the appended claims.

All references cited within the body of the instant specification are hereby incorporated by reference in their entirety.

5. EXAMPLES

5.1 EXAMPLE 1

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Novel Nucleic Acid Sequences Obtained From Various Libraries

A plurality of novel nucleic acids were obtained from cDNA libraries prepared from various human tissues and in some cases isolated from a genomic library derived from human chromosome using standard PCR, SBH sequence signature analysis and Sanger sequencing techniques. The inserts of the library were amplified with PCR using primers specific for the vector sequences which flank the inserts. Clones from cDNA libraries were spotted on nylon membrane filters and screened with oligonucleotide probes (e.g., 7-mers) to obtain signature sequences. The clones were clustered into groups of similar or identical sequences. Representative clones were selected for sequencing.

In some cases, the 5' sequence of the amplified inserts was then deduced using a typical Sanger sequencing protocol. PCR products were purified and subjected to fluorescent dye terminator cycle sequencing. Single pass gel sequencing was done using a 377 Applied Biosystems (ABI) sequencer to obtain the novel nucleic acid sequences

5.2 EXAMPLE 2

Assemblage of Novel Nucleic Acids

The nucleic acids of the present invention, designated as SEQ ID NO: 1-341 were assembled using an EST sequence as a seed. Then a recursive algorithm was used to extend the seed EST into an extended assemblage, by pulling additional sequences from different databases (i.e., Hyseq's database containing EST sequences, dbEST, gb pri, UniGene, and exons from public domain genomic sequences predicated by GenScan) that belong to this assemblage. The algorithm terminated when there was no additional sequences from the above databases that would extend the assemblage. Further, inclusion of component sequences into the assemblage was based on a BLASTN hit to the extending assemblage with BLAST score greater than 300 and percent identity greater than 95%.

Using PHRAP (Univ. of Washington) or CAP4 (Paracel), full-length gene sequences and their corresponding protein sequences were generated from the assemblage. Any frame shifts and incorrect stop codons were corrected by hand editing. During editing, the sequence was checked using FASTXY algorithm against Genbank (i.e., dbEST, gb pri, UniGene, and Genpept). Other computer programs which may have been used in the editing process were phredPhrap and Consed (University of Washington) and ed-ready, ed-ext and gc-zip-2 (Hyseq,

Inc.). The full-length nucleotide sequences are shown in the Sequence Listing as SEQ ID NO: 1-341. The corresponding polypeptide sequences are SEQ ID NO: 342-682.

Table 1 shows the various tissue sources of SEO ID NO: 1-341.

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The nearest neighbor results for polypeptides encoded by SEQ ID NO: 1-341 (i.e. SEQ ID NO: 342-682) were obtained by a BLASTP (version 2.0al 19MP-WashU) search against Genpept, Geneseq and SwissProt databases using BLAST algorithm. The nearest neighbor result showed the closest homologue with functional annotation for SEQ ID NO: 1-341. The translated amino acid sequences for which the nucleic acid sequence encodes are shown in the Sequence Listing. The homologues with identifiable functions for SEQ ID NO: 1-341 are shown in Table 2 below.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), polypeptides encoded by SEQ ID NO: 1-341 (i.e. SEQ ID NO: 342-682) were examined to determine whether they had identifiable signature regions. Table 3 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the Pfam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) polypeptides encoded by SEQ ID NO: 1-341 (i.e. SEQ ID NO: 342-682) were examined for domains with homology to certain peptide domains. Table 4 shows the name of the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

The GeneAtlas™ software package (Molecular Simulations Inc. (MSI), San Diego, CA) was used to predict the three-dimensional structure models for the polypeptides encoded by SEQ ID NO: 1-341 (i.e. SEQ ID NO: 342-682). Models were generated by (1) PSI-BLAST which is a multiple alignment sequence profile-based searching developed by Altschul et al, (Nucl. Acids. Res. 25, 3389-3408 (1997)), (2) High Throughput Modeling (HTM) (Molecular Simulations Inc. (MSI) San Diego, CA,) which is an automated sequence and structure searching procedure (http://www.msi.com/), and (3) SeqFold™ which is a fold recognition method described by Fischer and Eisenberg (J. Mol. Biol. 209, 779-791 (1998)). This analysis was carried out, in part, by comparing the polypeptides of the invention with the known NMR (nuclear magnetic resonance) and x-ray crystal three-dimensional structures

as templates. Table 5 shows, "PDB ID", the Protein DataBase (PDB) identifier given to

"Compound Information", information of the PDB template structure and/or its subcomponents; "PDB Function Annotation" gives function of the PDB template as annotated by the PDB files (http://www.rcsb.org/PDB/); start and end amino acid position of the protein sequence aligned; PSI-BLAST score, the verify score, the SeqFold score, and the Potential(s) of Mean Force (PMF). The verify score is produced by GeneAtlas™ software (MSI), is based on Dr. Eisenberg's Profile-3D threading program developed in Dr. David Eisenberg's laboratory (US patent no. 5,436,850 and Luthy, Bowie, and Eisenberg, Nature, 356:83-85 (1992)) and a publication by R. Sanchez and A. Sali, Proc. Natl. Acad. Sci. USA, 95:13597-12502. The verify score produced by GeneAtlas normalizes the verify score for proteins with different lengths so that a unified cutoff can be used to select good models as follows:

Verify score (normalized) = (raw score - 1/2 high score)/(1/2 high score)

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The PFM score, produced by GeneAtlas[™] software (MSI), is a composite scoring function that depends in part on the compactness of the model, sequence identity in the alignment used to build the model, pairwise and surface mean force potentials (MFP). As given in Table 5, a verify score between 0 to 1.0, with 1 being the best, represents a good model. Similarly, a PMF score between 0 to 1.0, with 1 being the best, represents a good model. A SeqFold[™] score of more than 50 is considered significant. A good model may also be determined by one of skill in the art based all the information in Table 5 taken in totality.

The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determined from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et al, as reference, were obtained for the polypeptide sequences. Table 6 shows the position of the last amino acid of the signal peptide in each of the polypeptides and the maximum score and mean score associated with that signal peptide.

Table 7 correlates each of SEQ ID NO: 1-341 to a specific chromosomal location.

Table 8 is a correlation table of the novel polynucleotide sequences SEQ ID NO: 1-341, and their corresponding priority nucleotide sequences in the priority application USSN 09/714,936, herein incorporated by reference in its entirety.

5

TABLE 1

Tissue	RNA	Library	SEQ ID NO:
Origin adult brain	Source GIBCO	Name AB3001	2 13 26-27 70 75 85 97 99-100 123 154-155 187-189
adult brain	GIBCO	ABD003	4 11 21 26-28 32 41 45 50 57 60-62 69-71 79 85 93 97 101 103-104
	1 02000	1.122005	113 115 117 126 131 142 150 154-155 177-178 181 184 190-201 225-
]		226 234 237 243 255-256
adult brain	Clontech	ABR001	6-7 11 14 26-27 75 93 107 131 154 201-202 243
adult brain	Clontech	ABR006	9 12 15 26-27 37 45 49 62 69 71 75 87 91 108-109 116 136 154 194
·	L	L	202 209 218-219 225 241 253 259 269-270 332 339
adult brain	Clontech	ABR008	2 6-7 9 12 15 18-22 26-28 35 37 40-41 45 48 50 55-56 61 63 65 67 71-
	1	ļ	76 78 85 91 94 99-101 105 108-109 117 121-123 130 140-142 145-
			147 149-152 154 158-159 170-174 185-186 189 198-199 201-202 205-
	}		206 212-213 220 225 228-229 236-237 240-242 248 252 255 259-262 269 272 281-282 286-287 297 302 318 326-327 339
adult brain	Clontech	ABR011	144 287
adult brain	BioChain	ABR012	23 232
adult brain	BioChain	ABR013	162
adult brain	Invitrogen	ABR014	37 40 87 253
adult brain	Invitrogen	ABR015	14 25 61 148
adult brain	Invitrogen	ABR016	40 61 124 126 225
adult brain	Invitrogen	ABT004	5 11 14-15 20 62 65 87 93-94 100 121 147 165 167 170 184-185 196
cultured	Church	400000	202 210 213 237 239-240 270 320
preadipocytes	Stratagene	ADP001	9 14 32 61 85 108-109 118 150 173 175-176 203 225
adrenal gland	Clontech	ADR002	11 13-14 18 21 33 43 64-65 99 101-102 104-106 108-109 111 126 156
actorial Piana	Cionicon	/IDROOZ	168 178 195 199 204 206 211 234 258 287
adult heart	GIBCO	AHR001	2 4 12 14-17 22 25 32-33 37 40-41 45 47-48 50 61 63-64 73-74 78 83
]	85 95 99 101 108-109 118 120 123-127 131 142 147 151-154 170 174
			203 212 225 227-228 236 244 249 259-260 271 287
adult kidney	GIBCO	AKD001	2 4-7 9 11-12 14-15 20-25 34 40-41 47-50 53 56 60-62 65 69-72 74
	ļ		76-79 83 85 87 90 93 95 97 99-100 103 108-110 113 116 118 121 123
			126-129 131 140 142 145-146 155-156 162 167 193 223 225 250-251 255 287
adult kidney	Invitrogen	AKT002	4-7 9 11 14 18 21 24-25 40 42-43 53 62 73 77 79 95 110 131 151-152
,			158 168 185 204 211 219 222 224 245 250-251 312
adult lung	GIBCO	ALG001	5 17 25-27 34 41 65 78 85 91 97 99 104 126 135 154 175 182 211 225
			233 330-331
lymph node	Clontech	ALN001	4 21 25-27 66 69 107 114 139 145-146 155 157 205 225 229
young liver	GIBCO	ALV001	4 10 12 14 24 40 59 64 94 100 103 105 121 139 154 198 234
adult liver	Invitrogen	ALV002	8 10 12 21 23 45 60 62-63 71 88 103 118 125 127 145-147 168 180
adult liver	Clontech	ALV003	198 224 257 266 303 322-323 266 337
adult ovary	Invitrogen	AOV001	2 4-7 9 11 13-16 18 21-23 25-27 33 35 37 40-41 43 45 47 52 57 60-65
	l mymogen	110 / 001	67 70-71 73 78-79 82 85 87-88 90-93 95 97-99 102 104-105 111 113-
			114 116-118 123 126-129 131 135 142 144-147 149-153 155 159-160
			164 166-172 174-175 177-179 182 185-186 190-194 196-197 206-209
			219 222 225 234-237 245-248 250-254 269-270 287 296 330-331
adult placenta	Invitrogen	APL001	20 37 61 69 216
placenta	Invitrogen	APL002	32 37 46 57 62 90 149 209
adult spleen	GIBCO	ASP001	4 14 20 25 32 41 45 49 61 68 70 78 93 97 99-100 103 118 131 138 142
adult testis	GIBCO	ATS001	148 151-152 158 162 175 177 201 216 222 225 234 309 2 11 14-15 20 35 40 61 76 81 97 113 127 145-146 159 200-201 206
addit testis		7110001	225 230 287
adult bladder	Invitrogen	BLD001	20 46 48 61-62 110 150 207 227 298
bone marrow	Clontech	BMD001	4 9 12 15 20 22 25-27 29 33 40-41 50-66 69-70 72 78 80-85 88 92 97
			102 108-109 113 115-116 120-121 130 132 141 148 162 178 191-192

Tissue Origin	RNA Source	Library Name	SEQ ID NO:
Origin	Source	Name	220 222 225 287 302
bone marrow	GF	BMD002	2 4 9 12 14-15 20-23 25-27 34-35 41-43 45 48 55-56 61-62 66 71 95 105-106 108-109 112 115-116 118 120 127 131 134 136 140-141 145-146 149 153 157 160 162 171-173 186 197 204 218 225 227 232 237 259-260 267 277 284 291 300 304 309 319 321 332 335 338
bone marrow	Clontech	BMD004	51
adult colon	Invitrogen	CLN001	13 21 87 93 97 130 140 149-150 164 199 232 250-251 266
mixture of 16 tissues/mRN As	various vendors	CTL021	16 61 213 225
mixture of 16 tissues/mRN As	various vendors	CTL028	61 216
adult cervix	BioChain	CVX001	2 5 14 17-18 21 32-33 40 42-43 50 61-62 64-65 70 74 78-79 82 89 92 95 97 110 114 123-124 127 155 158 168 170-172 175-177 185 197 224 234 250-251 265 287-289 333
endothelial cells	Stratagene	EDT001	2 4 11-16 18 20-21 23 26-27 32 34-35 40 42-44 47 49-50 56-57 61-63 65 70 72-74 85 88-91 93 95 99-100 106 108-110 117-118 123-124 126-129 142-143 145-146 160 175-178 190 194 204 206 209 216 225 236 262 287
Genomic clones from the short arm of chromosome 8	Genomic DNA from Genetic Research	EPM001	209
Genomic clones from the short arm of chromosome 8	Genomic DNA from Genetic Research	EPM003	209
Genomic clones from the short arm of chromosome 8	Genomic DNA from Genetic Research	EPM004	209
fetal brain	Clontech	FBR001	21 213
fetal brain	Clontech	FBR004	299
fetal brain	Clontech	FBR006	4 6-7 9 12 15 18-19 21 28-29 35 37 40 50 62 67 76 78 91 99 108-109 112 117 141 149 151-152 154 157 159 177 185 196 201-202 204 212 218 225 241 255 259 271 281 287 290 299-300 313 332 339
fetal brain	Invitrogen	FBT002	11-12 14 56 62 74 91 96 127 149 160 178-179 184-185 193 206 214 225 237 241-243
fetal heart	Invitrogen	FHR001	5 14 21 28 35 64-66 78 101 106 113 149 151-152 158 160 162 186 204 218 229 248 311 330-331 339-340
fetal kidney	Clontech	FKD001	12 23 33 40 61 69 82 91 98 104 155 175
fetal kidney	Clontech	FKD002	151-152 204 206 218 224 248 287
fetal kidney	Invitrogen	FKD007	25 61
fetal lung	Clontech	FLG001	21 35 126 159 203
fetal lung	Invitrogen	FLG003	6-7 14 23 45 48 56 61 121 149 154 164 180 234 248 250-251 330-331
fetal liver- spleen	Columbia University	FLS001	1-14 16-25 28-49 55 57 59 61-65 74 77-78 80 87-91 93-108 110-112 114 117-118 120-121 128-129 131 136 142-143 149 151-153 155 162 180-182 186 193 196 207 210-211 213 217-219 222 224 248 284 287 294 304 316 322

Tissue	RNA	Library	SEQ ID NO:
Origin	Source	Name	
fetal liver-	Columbia	FLS002	3-5 8 10 12-13 17 20-21 23-27 30-33 35-37 39-40 44 57 59 63-65 71-
spleen	University	1	72 74 77 79 88-89 93-95 97 99 101 103-107 111 114-115 117-118
			121-122 127-129 131 142 149 158 160 173 175-176 178 181-182 185
)		191-193 196 206-207 209-210 216-220 229 236 243 245-246 248-249
	ļ. <u></u>		257 277 294-296 311 317-318 325 341
fetal liver-	Columbia	FLS003	14 20 126 160 249 294 319 334
spleen	University	TT 7/001	671010111160100071070101101711601701701701
fetal liver	Invitrogen Clontech	FLV001	6-7 10 12 14 16 24 33 37 48 50 143 149 151-152 158 186 196 224 238
fetal liver	Clontech	FLV002 FLV004	14 21 61 149 335
Ictal livel	Ciontecn	FLV004	10 14 21 24 29 34-35 37 45 47 69 72 108-109 116 118 139 157 179 255 332
fetal muscle	Invitrogen	FMS001	21 26-27 32 35 37 44 61 94 108-109 118 124 126-127 134 159 190
	l	11.13001	216 263
fetal muscle	Invitrogen	FMS002	14 21-22 42-43 67-68 85 108-109 111 118-119 145-146 185 198 216
		!	262-263 332 336 339
fetal skin	Invitrogen	FSK001	2 10-14 17 28 33 37 40 46 59 62-63 68-69 71 81 90 93 100 115 122
			127 131 143 150 153 156 160 174 195-196 206 213 216 224-225 239
			287 301-302 313-315
fetal skin	Invitrogen	FSK002	2 22 34 41 66 71 100 113-114 116 121 143 178-179 194 209 216 227
			259 267 313
fetal spleen	BioChain	FSP001	21 91
umbilical	BioChain	FUC001	2 14 17 21 25-27 33 42-43 45 48 60-62 78 85-86 90 93 97 99 103 107
fetal brain	GIBCO	UEDOOL	110 116-117 126 147 151-152 161 168 216 220 234 236 283
letai brain	GIBCO	HFB001	14-15 18 21 23 26-28 32 35 40-41 43 47 60 67-68 70-79 85 94 99 101 144-146 149 151-152 158 177 183-184 197 212-213 225
infant brain	Columbia	IB2002	4-5 9 11-12 14 16 21 28-29 35 37 47-48 64 68 71-72 75 79 91-93 99-
	University	152002	100 103 106 121 126 131 147 151-152 154-155 159 162 177 182 185-
			187 201 209 211 213-214 225 246 267 271 309 319-320 328
infant brain	Columbia	IB2003	4-5 9 21 26-28 45 79 90 92-93 131 147-148 185 191-192 205 213-214
	University		336
infant brain	Columbia	IBM002	21 75 320
	University		
infant brain	Columbia	IBS001	21 150 185 320
61-11-1	University	1 55 661	
fibroblast	Stratagene	LFB001	2 13-14 18 26-27 33 40 42-43 93 99 111 116 123 126 133 137 150 155
adult lung	Invitrogen	LGT002	175-176 201 216 225 245 329 5-7 11 14 20-21 26-27 33 35 37 40-43 47-48 53 59 61-62 72 74 79 81
addit fullg	mvinogen	LG1002	83 85 90-91 95 97 99-100 104 106-107 111 117-118 126-127 136 139-
			140 142 145-146 153 155 160 162 164 170 175-176 181-182 203 206
			215-216 220-225 233-235 248-251 262 268 291 309-310 330-331
lymphocytes	ATCC	LPC001	4 9 14 21 26-27 41 50 61 69 83 100 107 113 117-118 120 131 137 164
			170-172 209 225 227 245 247 275 286 319
leukocyte	GIBCO	LUC001	1-2 4-5 9 12-15 20-22 25-27 33 35 38 40-43 50 53 57 59-63 65 69 71-
			72 74 76 78-79 82-83 88 93 95 97-99 101 103 107-109 113-114 116-
			120 123 126 131 133-139 150 161-165 173 178 218 222 225 227 250-
Invites a d	01-4:1	TYTOOO	251 273-275 287 305-307 309 319 338
leukocyte	Clontech	LUC003	4-5 12 42-43 63 71 99 116 118 148 162 166 171-172 309
melanoma fro mcell line	Clontech	MEL004	2 9 12 20 26-27 70 72 79 100 113 116 126 147-148 168 184 218 225
ATCC #CRL		1	284 304
1424			
mammary	Invitrogen	MMG001	5-7 12-16 20-21 28 32 45-46 48 59 61-62 65 71 74 79 90-91 93-94 97
gland			100 102-103 110 115 118 121-122 131 139 149 162 167 169 196 198
			206-207 216 220 222 224-225 233 236 245 255-258 287 311 330-331
		-	339
induced	Stratagene	NTD001	13-14 26-27 32 61 65 72 78

Tissue	RNA	Library	SEQ ID NO:
Origin	Source	Name	
neuronal cells			
retinoic acid-	Stratagene	NTR001	14 16 44 231 249
induced			
neuronal cells			
neuronal cells	Stratagene	NTU001	5 13-14 16 21 68 72 74 115 150 160 170
pituitary	Clontech	PIT004	9 34 69 74 85 99 270 333
gland			
placenta	Clontech	PLA003	9 35 37 45 64 87 93 99 113 116 139 164 218
prostate	Clontech	PRT001	14 17 21-22 33-34 63-64 79 85 93 99 111 158 200 225 245 262 275
rectum	Invitrogen	REC001	5-7 13 20 41 61-63 93 100 110-111 130 149 158 199 206 218 223 245
			302-303 320
salivary gland	Clontech	SAL001	5 14 23 61 70 91 105 111 137 162 245 276 285
skin fibroblast	ATCC	SFB002	225
small	Clontech	SIN001	12 14 17 21-22 41 44 46-47 60 62 71-72 83 86 94 100 105 121 126
intestine	Į		131 136 138 171-173 175 183 185 203 205 207 216-217 233 235 238-
			239 245 250-251 276 285 335
skeletal	Clontech	SKM001	12 14 26-27 35 76 91 103 118 263
muscle			
spinal cord	Clontech	SPC001	5 14 21-22 25-27 34 45 48 61 67 70-71 76 91 95 118 126-127 154 173
			199 212 222 225 281
Adult spleen	Clontech	SPLc01	1 33 41 121 222
stomach	Clontech	STO001	4 21 25-27 38 53 61 63 76 104 111 115 155 215 225 238 262 275 277
thalamus	Clontech	THA002	20 37 74 111 114 130 149 187 193 206 209 216 250-251 253 261
thymus	Clontech	THM001	4 9 14 18 21 23 26-27 41-43 59 61 69 83 95 100 106 114 124 126 128-
			129 133 155 170 178 245 258 277
thymus	Clontech	THMc02	12 20-21 26-27 35 40 48 59 61-62 64 66 69 78 94 99 106-109 112-113
1			118 126-129 140 157 161-162 164 170 173 191-192 208-209 213 221-
			222 253 260 278 286 291-292 305 307-309 316
thyroid gland	Clontech	THR001	4-7 11-12 14-15 18 21-23 33 35 40 46 59 69-74 76 78 83 85-86 91 93
			97 105-106 108-109 114 117 123 126 131 138-139 145-146 151-153
			165 173 190 194 206 225 234 263 265 276 279-280 293 297 324
trachea	Clontech	TRC001	49-50 60 62 73-74 76 88 134 178 225 250-251 264-265
uterus	Clontech	UTR001	2 5 12 14 17 21 26-27 33 50 69 85 97 117 138

The 16 tissue/mRNAs and their vendor sources are as follows: 1) Normal adult brain mRNA (Invitrogen), 2) Normal adult kidney mRNA (Invitrogen), 3) Normal fetal brain mRNA (Invitrogen), 4) Normal adult liver mRNA (Invitrogen), 5) Normal fetal kidney mRNA (Invitrogen), 6) Normal fetal liver mRNA (Invitrogen), 7) normal fetal skin mRNA (Invitrogen), 8) human adrenal gland mRNA (Clontech), 9) Human bone marrow mRNA (Clontech), 10) Human leukemia lymphoblastic mRNA (Clontech), 11) Human thymus mRNA (Clontech), 12) human lymph node mRNA (Clontech), 13) human so\spinal cord mRNA (Clontech), 14) human thyroid mRNA (Clontech), 15) human esophagus mRNA (BioChain), 16) human conceptional umbilical cord mRNA (BioChain).

TABLE 2

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
342	AK027819	Homo sapiens	FLJ14913 fis, clone PLACE1006782.	2806	100
343	AAB81047	Homo sapiens	20-JUN-2001 28-JUL-1999 Human protein	1708	100
			HP00698 amino acid sequence.		
344	AB040926	Homo sapiens	for KIAA1493 protein, partial cds.	1973	98
345	AAB01382	Homo sapiens	20-OCT-2000 10-DEC-1999 Neuron- associated protein.	4363	99
346	AAY99410	Homo sapiens	08-AUG-2000 01-SEP-1999 Human PRO1480 (UNQ749) amino acid sequence SEQ ID NO:253.	3576	99
347	AAE01114	Homo sapiens	17-JUL-2001 08-NOV-2000 Human gene 1 encoded secreted protein HBINK72, SEQ ID NO:28.	2767	99
348	AAE01114	Homo sapiens	17-JUL-2001 08-NOV-2000 Human gene 1 encoded secreted protein HBINK72, SEQ ID NO:28.	1652	76
350	AF113208	Homo sapiens	mRNA, complete cds.	1615	100
351	AAB49535	Homo sapiens	09-MAR-2001 06-APR-2000 Clone HFKCD20.	3027	100
352	BC001079	Homo sapiens	clone MGC:2731 IMAGE:2822460, mRNA, complete cds.	1127	99
353	AAB20093	Homo sapiens	23-APR-2001 16-JUN-2000 Human hydrophobic domain-containing protein HP03374.	803	100
354	AY007148	Homo sapiens	CDABP0084 mRNA sequence.	984	100
355	BC001795	Homo sapiens	Similar to ribosomal protein S2, clone MGC:3141 IMAGE:3353508, mRNA, complete cds.	971	100
356 ·	BC008739	Homo sapiens	protein x 013, clone MGC:3073 IMAGE:3346340, mRNA, complete cds.	386	100
357	AY007133	Homo sapiens	CDABP0047 mRNA sequence.	1639	95
358	X15977	Homo sapiens	mRNA for collagen VI alpha-2 alternative C-terminal domain.	515	100
359	BC013173	Homo sapiens	clone MGC:17340 IMAGE:4340287, mRNA, complete cds.	3049	100
360	BC011747	Homo sapiens	Similar to secretory carrier membrane protein 4, clone MGC:19661 IMAGE:3161979, mRNA, complete cds.	1022	87
363	AJ310550	Homo sapiens	for SMC5 protein.	3517	99
364	AJ276485	Homo sapiens	for putative integral membrane transporter protein (LC27 gene).	1502	100
365	J05158	Homo sapiens	carboxypeptidase N mRNA, 3' end.	2274	88
366	X57351	Homo sapiens	1-8D gene from interferon-inducible gene family.	673	97
367	AF230904	Homo sapiens	protein (CIN85) mRNA, complete cds.	3437	100
368	AF230904	Homo sapiens	protein (CIN85) mRNA, complete cds.	2615	99
369	AJ236915	Homo sapiens	for pak5 protein.	3550	100
370	AF269255	Homo sapiens	apyrase-like protein 1 (LALP1) mRNA, complete cds.	3198	100
373	AAY24791	Homo sapiens	26-AUG-1999 18-DEC-1998 Human secreted protein nm134_4.	1277	100
374	X68277	Homo sapiens	CL 100 mRNA for protein tyrosine phosphatase.	1886	100
375	AK025844	Homo sapiens	FLJ22191 fis, clone HRC01066.	1904	100
376	AF032668	Rattus	rsec15	3738	92

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
		norvegicus			
378	AF195534	Rattus norvegicus	GERp95	4513	99
379	AAG63221	Homo sapiens	01-OCT-2001 18-JAN-2001 Amino acid sequence of a human lipid metabolism enzyme.	518	100
380	AAB68878	Homo sapiens	24-APR-2001 21-JUL-2000 Human RECAP polypeptide, SEQ ID NO: 8.	946	100
381	BC004546	Homo sapiens	disrupter of silencing 10, clone MGC:11290 IMAGE:3946633, mRNA, complete cds.	2431	100
382	AAY02361	Homo sapiens	13-JUL-1999 06-OCT-1998 Polypeptide identified by the signal sequence trap method.	979	98
383	AAB63460	Homo sapiens	26-MAR-2001 26-MAY-2000 Human breast cancer associated antigen protein sequence SEQ ID NO:822.	984	99
384	AAB63460	Homo sapiens	26-MAR-2001 26-MAY-2000 Human breast cancer associated antigen protein sequence SEQ ID NO:822.	984	99
385	BC001068	Homo sapiens	clone IMAGE:2823731, mRNA, partial cds.	2994	99
386	AK003950	Mus musculus	putative	623	97
387	AK001527	Homo sapiens	FLJ10665 fis, clone NT2RP2006200.	4109	99
388	BC014442	Homo sapiens	clone MGC:22964 IMAGE:4866321, mRNA, complete cds.	2333	100
389	BC000056	Homo sapiens	clone MGC:3262 IMAGE:3506385, mRNA, complete cds.	1464	95
390	BC004393	Homo sapiens	Similar to RIKEN cDNA 2310045B01 gene, clone MGC:10974 IMAGE:3635540, mRNA, complete cds.	1145	99
391	AK026302	Homo sapiens	FLJ22649 fis, clone HSI07332.	930	99
392	AK001411	Homo sapiens	FLJ10549 fis, clone NT2RP2001976, moderately similar to Mus musculus	3711	100
393	AAB93202	Homo sapiens	calmodulin-binding protein SHA1 mRNA. 26-JUN-2001 28-JUL-2000 Human protein sequence SEQ ID NO:12168.	2549	99
394	AAG75102	Homo sapiens	03-SEP-2001 28-SEP-2000 Human colon cancer antigen protein SEQ ID NO:5866.	995	100
396	AF006088	Homo sapiens	protein complex subunit p16-Arc (ARC16) mRNA, complete cds.	371	100
397	BC005131	Homo sapiens	Similar to RIKEN cDNA 2010003J03 gene, clone MGC:11102 IMAGE:3831647, mRNA, complete cds.	849	99
398	AK010289	Mus musculus	putative	854	73
399	AF226055	Homo sapiens	(HTGN29) mRNA, complete cds.	1367	100
400	AF090930	Homo sapiens	HQ0478 PRO0478 mRNA, complete cds.	180	89
401	AF118084	Homo sapiens	PRO1914	350	98
402	BC007283	Homo sapiens	ribosomal protein S11, clone MGC:15628 IMAGE:3343839, mRNA, complete cds.	824	100
403	AK025392	Homo sapiens	FLJ21739 fis, clone COLF4061.	4331	99
404	AF077615	Homo sapiens	beta inducible nuclear protein TINP1 (TINP1) mRNA, complete cds.	1364	100
405	AK027709	Homo sapiens	FLJ14803 fis, clone NT2RP4001442.	2963	99
406	BC006002	Homo sapiens	Similar to RIKEN cDNA 1190005P17 gene, clone MGC:14817 IMAGE:4247279, mRNA, complete cds.	666	100
407	M80902	Homo sapiens	AHNAK nucleoprotein mRNA, 5' end.	8529	99
408	AAW90962	Homo sapiens	14-JUL-2000 06-NOV-1998 Human CSGP-2	2346	99

SEQ ID	Accession	Species	Description	Score	%
NO:	No.				Identity
400	17500000		protein.		
409	AK027715	Homo sapiens	FLJ14809 fis, clone NT2RP4001822, weakly similar to PLATELET-ENDOTHELIAL TETRASPAN ANTIGEN 3.	1295	100
410	BC015928	Homo sapiens	clone MGC:8773 IMAGE:3908916, mRNA, complete cds.	2186	100
411	BC015317	Homo sapiens	Similar to suppression of tumorigenicity 13 (colon carcinoma) (Hsp70-interacting protein), clone MGC:21083 IMAGE:4425762, mRNA, complete cds.	302	100
412	L26335	Cavia porcellus	zinc finger protein	1493	99
413	AF209198	Homo sapiens	finger protein 277 (ZNF277) mRNA, complete cds.	2357	100
414	AE001399	Plasmodium falciparum	GAF domain protein (cyclic nt signal transduct.)	178	35
415	AAY48226	Homo sapiens	08-DEC-1999 10-MAR-1998 Human prostate cancer-associated protein 12.	1204	96
416	M94389	Loligo pealei	neurofilament protein	165	23
417	AF317425	Homo sapiens	(GAC-1) mRNA, complete cds.	3725	91
418	AF116675	Homo sapiens	PRO1942	257	100
419	AAG73932	Homo sapiens	03-SEP-2001 28-SEP-2000 Human colon cancer antigen protein SEQ ID NO:4696.	1415	100
420	AK000100	Homo sapiens	FLJ20093 fis, clone COL04263.	841	100
421	BC005326	Homo sapiens	ribosomal protein L27a, clone MGC:12412 IMAGE:4052417, mRNA, complete cds.	754	99
422	AF119865	Homo sapiens	PRO2176	470	97
424	AF138863	Homo sapiens	PRO1677	868	99
425	X14361	Homo sapiens	CR1 gene for C3b/C4b receptor SCR9 (or 16) C-term. exon SCR = short consensus repeat.	135	100
426	Z24725	Homo sapiens	mitogen inducible gene mig-2, complete CDS.	3576	99
427	AK027587	Homo sapiens	FLJ14681 fis, clone NT2RP2004270, weakly similar to PROTEIN PTM1 PRECURSOR.	1103	100
428	AC004770	Homo sapiens	11, BAC CIT-HSP-311e8 (BC269730) containing the hFEN1 gene, complete sequence.	1527	84
429	AK026262	Homo sapiens	FLJ22609 fis, clone HSI04913.	1795	99
430	BC007279	Homo sapiens	clone FLB5214, clone MGC:15622 IMAGE:3343280, mRNA, complete cds.	416	100
431	AL133035	Homo sapiens	cDNA DKFZp434G171 (from clone DKFZp434G171).	1136	99
432	AF166125	Homo sapiens	N mRNA, partial cds.	1816	99
433	AF161370	Homo sapiens	mRNA, partial cds.	824	100
434	AK000161	Homo sapiens	FLJ20154 fis, clone COL08740.	284	100
435	AK001784	Homo sapiens	FLJ10922 fis, clone OVARC1000420.	684	100
436	BC011396	Homo sapiens	clone MGC:17720 IMAGE:3870711, mRNA, complete cds.	1080	100
437	AF165527	Homo sapiens	(DGCR8) mRNA, complete cds.	859	100
438	AF230200	Homo sapiens	mRNA, partial cds.	358	95
439	BC008468	Homo sapiens	Similar to RIKEN cDNA 1110059G10 gene, clone MGC:14734 IMAGE:4277104, mRNA, complete cds.	791	100
440	BC007870	Homo sapiens	DC6 protein, clone MGC:14435 IMAGE:4303290, mRNA, complete cds.	505	100
441	AAB20167	Homo sapiens	30-APR-2001 17-JUL-2000 Human protein	2066	100

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
			associated with IgA nephropathy.		
442	AAB08910	Homo sapiens	30-AUG-2000 22-SEP-1999 Human secreted protein sequence encoded by gene 20 SEQ ID NO:67.	1112	100
443	BC003026	Homo sapiens	clone IMAGE:2823490, mRNA, partial cds.	354	84
444	BC003127	Homo sapiens	Similar to selenoprotein X, 1, clone MGC:3344 IMAGE:2905838, mRNA, complete cds.	527	100
445	AK000143	Homo sapiens	FLJ20136 fis, clone COL07068.	2260	100
446	AK000388	Homo sapiens	FLJ20381 fis, clone KAlA2329.	2375	100
447	BC002364	Homo sapiens	non-POU-domain-containing, octamer- binding, clone MGC:8677 IMAGE:2964534, mRNA, complete cds.	2449	98
448	AK025645	Homo sapiens	FLJ21992 fis, clone HEP06554.	920	88
449	AAB95268	Homo sapiens	26-JUN-2001 28-JUL-2000 Human protein sequence SEQ ID NO:17462.	3708	99
450	AF113538	Homo sapiens	x receptor interacting protein mRNA, complete cds.	1800	100
451	AAW78167	Homo sapiens	13-APR-1999 11-JUN-1998 Human secreted protein encoded by gene 42 clone HFFAT33.	795	100
452	BC014943	Homo sapiens	NMN adenylyltransferase; nicotinamide mononucleotide adenylyl transferase, clone MGC:22925 IMAGE:4874147, mRNA, complete cds.	1458	100
453	BC000348	Homo sapiens	ribosomal protein L35, clone MGC:8582 IMAGE:2960987, mRNA, complete cds.	591	97
454	AJ277591.	Homo sapiens	for p15-2a protein (p15-2 gene).	749	100
455	AK000927	Homo sapiens	FLJ10065 fis, clone HEMBA1001455.	3143	100
456	AB045118	Homo sapiens	mRNA, complete cds.	1192	99
457	AAZ51355	Homo sapiens	06-JUN-2000 20-AUG-1999 Human wild type serine/threonine kinase KIS (hKIS) gene.	2198	99
458	AF146696	Homo sapiens	pAB195 FOXP1 (FOXP1) mRNA, complete cds.	1639	100
459	BC009401	Homo sapiens	natural killer cell transcript 4, clone MGC:15353 IMAGE:4300407, mRNA, complete cds.	914	100
460	BC010537	Homo sapiens	activated RNA polymerase II transcription cofactor 4, clone MGC:17295 IMAGE:3457167, mRNA, complete cds.	563	99
461	AF076642	Homo sapiens	of G-protein signaling 13 mRNA, complete cds.	1218	100
462	AF116718	Homo sapiens	PRO2900	396	100
463	AAB18919	Homo sapiens	08-FEB-2001 01-MAR-2000 A novel polypeptide designated PRO4356.	1137	99
464	AC025416	Arabidopsis thaliana	F5O11.12	135	36
465	BC002757	Homo sapiens	cytochrome c oxidase subunit VIIa polypeptide 1 (muscle), clone MGC:3716 IMAGE:3631740, mRNA, complete cds.	247	100
466	AY037115	Homo sapiens	stromal lymphopoietin (TSLP) mRNA, complete cds.	823	100
467	M15841	Homo sapiens	U2 small nuclear RNA-associated B" antigen mRNA, complete cds.	638	100
468	AK026916	Homo sapiens	FLJ23263 fis, clone COL06129.	2612	99
469	AAY05317	Homo sapiens	25-JUN-1999 08-SEP-1998 Human secreted protein bn97_1.	1508	100

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
470	AAY05317	Homo sapiens	25-JUN-1999 08-SEP-1998 Human secreted protein bn97_1.	851	99
471	AAY66721	Homo sapiens	05-APR-2000 02-JUN-1999 Membrane- bound protein PRO511.	1176	95
472	AAB12144	Homo sapiens	02-FEB-2001 17-NOV-1999 Hydrophobic domain protein isolated from WERI-RB cells.	1806	100
474	AL022398	Homo sapiens	sequence from PAC 434014 on chromosome 1q32.341. Contains the HSD11B1 gene for Hydroxysteroid (11-beta) Dehydrogenase 1, the ADORA2BP adenosine A2b receptor LIKE pseudogene, the IRF6 gene for Interferon Regulatory Factor 6 and two novel genes. Contains ESTs and GSSs, complete sequence.	575	100
475	AF324830	Homo sapiens	transcript 11 protein (ILT11) mRNA, complete cds.	1590	100
476	AJ306731	Homo sapiens	for RhoGAP protein (RICH1 gene).	846	100
477	BC006116	Homo sapiens	Similar to RIKEN cDNA 3100002B05 gene, clone MGC:12993 IMAGE:3504453, mRNA, complete cds.	2063	100
478	AK001077	Homo sapiens	FLJ10215 fis, clone HEMBA1006737, weakly similar to ANKYRIN, BRAIN VARIANT 2.	812	100
479	AAG89322	Homo sapiens	11-SEP-2001 07-DEC-2000 Human secreted protein, SEQ ID NO: 442.	922	98
480	AAE02782	Homo sapiens	06-AUG-2001 06-DEC-2000 Human six transmembrane epithelial antigen of prostate (STEAP)-3 protein.	2392	100
481	AK025537	Homo sapiens	FLJ21884 fis, clone HEP02863.	3021	99
482	AJ007590	Homo sapiens	for XRP2 protein.	1766	100
483	AAG93264	Homo sapiens	13-SEP-2001 06-DEC-2000 Human protein HP10160.	841	100
484	AB027258	Homo sapiens	for basal transcriptional activator hABT1, complete cds.	1408	100
485	BC000518	Homo sapiens	Similar to brain acid-soluble protein 1, clone MGC:8555 IMAGE:2822874, mRNA, complete cds.	1137	99
486	AK001425	Homo sapiens	FLJ10563 fis, clone NT2RP2002769.	1695	99
487	BC013322	Homo sapiens	clone MGC:13411 IMAGE:4077631, mRNA, complete cds.	1459	99
488	AK002030	Homo sapiens	FLJ11168 fis, clone PLACE1007274.	1029	100
489	BC003378	Homo sapiens	high-mobility group (nonhistone chromosomal) protein 1, clone MGC:5223 IMAGE:2901382, mRNA, complete cds.	1140	99
490	AK001159	Homo sapiens	FLJ10297 fis, clone NT2RM1001074.	764	100
491	AK000020	Homo sapiens	FLJ20013 fis, clone ADKA03455.	1613	100
492	AK001322	Homo sapiens	FLJ10460 fis, clone NT2RP1001475.	1207	100
493	AK001322	Homo sapiens	FLJ10460 fis, clone NT2RP1001475.	892	98
494	AY008293	Homo sapiens	protease (SENP8) mRNA, complete cds.	1114	99
495	AF413080	Homo sapiens	mRNA, complete cds.	9184	99
496	AK000154	Homo sapiens	FLJ20147 fis, clone COL07954.	673	100
497	AK001001	Homo sapiens	FLJ10139 fis, clone HEMBA1003175.	658	100
499 501	AK027124	Homo sapiens	FLJ23471 fis, clone HSI11969.	1773	99
201	BC012024	Homo sapiens	kinetochore protein CENP-H, clone MGC:21431 IMAGE:4510607, mRNA, complete cds.	1214	100

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
502	U40407	synthetic construct	T cell receptor alpha chain	1119	80
503	AF043179	Homo sapiens	cell receptor beta chain (TCRBV13S1- TCRBJ2S1) mRNA, complete cds.	681	73
504	AF116678	Homo sapiens	PRO1995	587	100
505	AB051853	Homo sapiens	gene for rho-GTPase activating protein, complete cds.	1766	98
506	AB046074	Macaca fascicularis	unnamed protein product	515	83
507	AK002848	Mus musculus	putative	429	84
508	AAB08973	Homo sapiens	30-AUG-2000 22-SEP-1999 Human secreted protein sequence encoded by gene 27 SEQ ID NO:130.	1753	98
509	AK000740	Homo sapiens	FLJ20733 fis, clone HEP08550.	4651	100
510	AL136858	Homo sapiens	cDNA DKFZp434N2435 (from clone DKFZp434N2435); complete cds.	501	100
511	BC008413	Homo sapiens	clone MGC:14552 IMAGE:4333393, mRNA, complete cds.	1706	99
513	AJ277275	Homo sapiens	for rapa-1 (rapa gene).	5086	100
514	AB042563	Homo sapiens	mRNA for casein kinase 1 gamma 1L, complete cds.	1739	100
515	BC015597	Homo sapiens	clone IMAGE:4649498, mRNA, partial cds.	719	63
516	BC001277	Homo sapiens	KDEL (Lys-Asp-Glu-Leu) endoplasmic reticulum protein retention receptor 3, clone MGC:5099 IMAGE:3462392, mRNA, complete cds.	1103	100
517	AF081126	Drosophila melanogaster	ER lumen protein retaining receptor	409	75
519	AK023651	Homo sapiens	FLJ13589 fis, clone PLACE1009308, weakly similar to GLUCOSE REPRESSION MEDIATOR PROTEIN.	1488	100
520	AK000371	Homo sapiens	FLJ20364 fis, clone HEP17854.	2040	100
522	AAB24228	Homo sapiens	07-FEB-2001 06-APR-2000 Human vesicle associated protein 7 SEQ ID NO:7.	1293	100
523	BC015387	Homo sapiens	Similar to RIKEN cDNA 1110001019 gene, clone MGC:21689 IMAGE:4400374, mRNA, complete cds.	429	100
524	BC008488	Homo sapiens	RIKEN cDNA 2010100012 gene, clone MGC:14813 IMAGE:4133274, mRNA, complete cds.	404	97
526	AF360739	Homo sapiens	protein SS-56 (SS-56) mRNA, complete cds.	2618	99
527	BC015725	Homo sapiens	clone MGC:17998 IMAGE:3922049, mRNA, complete cds.	782	100
529	AF230201	Homo sapiens	mRNA, complete cds.	396	100
530	AK001984	Homo sapiens	FLJ11122 fis, clone PLACE1006159.	658	100
531	AK000530	Homo sapiens	FLJ20523 fis, clone KAT10456.	691	100
532	U37134	Drosophila melanogaster	inturned protein	248	23
533	U37134	Drosophila melanogaster	inturned protein	244	23
535	AB033132	Homo sapiens	complete cds, testis-specific gene2.	1586	100
536	AF153417	Homo sapiens	9 open reading frame 6 mRNA, complete cds.	221	100
537	AJ277557	Homo sapiens	gene for mitochondrial 5'(3')- deoxyribonucleotidase (dNT-2 gene), exons 1-5.	617	100
538	AF127564	Arabidopsis	ubiquitin-protein ligase 1	854	42
			110		

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
		thaliana			ruchity
540	AK000442	Homo sapiens	FLJ20435 fis, clone KAT03864.	1513	99
541	AF278541	Homo sapiens	protein ACT mRNA, complete cds.	1657	99
542	AAY99440	Homo sapiens	08-AUG-2000 01-SEP-1999 Human	3408	100
			PRO1564 (UNQ770) amino acid sequence	5.00	'''
	Ĭ		SEQ ID NO:347.	1	
543	AL117491	Homo sapiens	cDNA DKFZp434N231 (from clone	7295	100
		•	DKFZp434N231); partial cds.		
544	BC003179	Homo sapiens	clone MGC:4419 IMAGE:2958058, mRNA,	792	100
			complete cds.		
545	AAE05186	Homo sapiens	12-SEP-2001 12-JAN-2001 Human drug	1095	99
			metabolising enzyme (DME-17) protein.		
546	AAY94926	Homo sapiens	16-JUN-2000 13-AUG-1999 Human secreted	1578	99
			protein clone rd232_5 protein sequence SEQ		
		<u> </u>	ID NO:58.		
547	AK026027	Homo sapiens	FLJ22374 fis, clone HRC06766.	647	100
548	AL137584	Homo sapiens	cDNA DKFZp434G1310 (from clone	246	97
550	1 D0 5000 6		DKFZp434G1310); partial cds.		
550	AF352026	Homo sapiens	protein 1 mRNA, complete cds.	3085	99
552	AK025840	Homo sapiens	FLJ22187 fis, clone HRC01029.	918	100
553	BC013117	Homo sapiens	clone MGC:8711 IMAGE:3882749, mRNA,	1126	100
554	BC014111	III	complete cds.	0.000	\
334	BC014111	Homo sapiens	Similar to ecotropic viral integration site 5,	2698	97
	•		clone MGC:20844 IMAGE:4542709, mRNA, complete cds.	}	
555	AK016622	Mus musculus	putative	1412	07
557	AF181263	Homo sapiens	domain containing 2 (EHD2) mRNA,	1413 2816	97 99
337	A1 101203	110mo sapiciis	complete cds.	2010	1 99
558	AP001660	Homo sapiens	DNA, chromosome 21q, section 4/105.	1424	100
559	BC001781	Homo sapiens	ribosomal protein L44, clone MGC:2064	542	100
		axomo supremo	IMAGE:3353669, mRNA, complete cds.	342	100
560	AF081941	Rattus	soluble adenylyl cyclase	142	38
		norvegicus			
561	AF378129	Homo sapiens	domain containing adaptor protein TIRAP	1227	99
		·	mRNA, complete cds.		
562	X01403	Homo sapiens	mRNA fragment for T-cell receptor alpha	840	90
			chain.		
563	AAY39883	Homo sapiens	07-DEC-1999 26-MAR-1999 MHC Class II	947	99
			p41 specific region.		
564	AB026707	Homo sapiens	for FOAP-11 protein, complete cds.	429	100
565	AK007905	Mus musculus	putative	1484	83
566	BC015389	Homo sapiens	clone IMAGE:4401937, mRNA, partial cds.	421	100
567	AF116669	Homo sapiens	PRO1828	237	100
568	AK000328	Homo sapiens	FLJ20321 fis, clone HEP09380.	5507	99
569	AF263913	Mus musculus	fidgetin	3864	97
570	AK015017	Mus musculus	putative	635	50
572	AK001673	Homo sapiens	FLJ10811 fis, clone NT2RP4000955.	3661	100
573	AAY96059	Homo sapiens	05-DEC-2000 02-MAR-2000 Human	617	100
	177000000	 , , 	sphingosine kinase C.		
574	AK000207	Homo sapiens	FLJ20200 fis, clone COLF1206.	2500	99
575	X52140	Rattus	precursor polypeptide (AA -28 to 1152)	5429	87
576	A 1/20050000	norvegicus			
					100
3/1	AABU88/U	riomo sapiens		590	100
576 577	AK005909 AAB08870	Mus musculus Homo sapiens	putative 15-JAN-2001 03-MAR-2000 Amino acid sequence of a human secretory protein.	393 590	

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
578	AJ296173	Mus musculus	GATS protein	582	96
580	AE003588	Drosophila melanogaster	CG13947 gene product	115	42
582	AK023117	Homo sapiens	FLJ13055 fis, clone NT2RP3001538, weakly similar to HYPOTHETICAL 39.0 KD PROTEIN T28D9.3 IN CHROMOSOME II.	1664	99
583	BC011870	Homo sapiens	Similar to mesenchymal stem cell protein DSC43, clone MGC:19952 IMAGE:2960099, mRNA, complete cds.	1554	100
585	BC003563	Homo sapiens	guanine nucleotide binding protein (G protein), gamma 5, clone MGC:1969 IMAGE:3502879, mRNA, complete cds.	333	98
586	AL035521	Arabidopsis thaliana	putative protein	145	28
587	AY014283	Homo sapiens	mRNA, complete cds.	1066	100
588	AK020796	Mus musculus	putative	519	85
589	AL034548	Homo sapiens	DNA sequence from clone RP5-1103G7 on chromosome 20p12.2-13. Contains up to three novel genes, the gene for a novel protein similar to mouse VMP, the gene for a novel protein kinase domains containing protein similar to phosphoprotein C8FW and rat NIPK, and the SOX22 gene for SRY (sexdetermining region Y)-box 22. Contains five CpG islands, ESTs, STSs and GSSs, complete sequence.	262	100
590	AK023084	Homo sapiens	FLJ13022 fis, clone NT2RP3000753, weakly similar to NEUROFILAMENT TRIPLET H PROTEIN.	1144	99
591	X97966	Homo sapiens	mRNA for calcyphosine.	963	100
592	X97966	Homo sapiens	mRNA for calcyphosine.	660	95
594	BC002471	Homo sapiens	complexin 1, clone MGC:3097 IMAGE:3349779, mRNA, complete cds.	668	99 .
596	BC007394	Homo sapiens	clone MGC:16291 IMAGE:3834089, mRNA, complete cds.	217	85
598	X85738	Bos taurus	novel brain-specific protein	326	55
600	AJ310550	Homo sapiens	for SMC5 protein.	880	97
601	BC001466	Homo sapiens	ring-box 1, clone MGC:1481 IMAGE:3138751, mRNA, complete cds.	131	100
602	AK012283	Mus musculus	putative	1711	96
603 605	AF251062 AAG02234	Homo sapiens Homo sapiens	binding protein mRNA, complete cds. 06-OCT-2000 21-FEB-2000 Human secreted	1551 284	99
			protein, SEQ ID NO: 6315.		
606	AAG01931	Homo sapiens	06-OCT-2000 21-FEB-2000 Human secreted protein, SEQ ID NO: 6012.	159	73
608	AK001757	Homo sapiens	FLJ10895 fis, clone NT2RP4002905.	1300	100
610	U20897	Homo sapiens	clone 475/1 melanoma ubiquitous mutated protein (MUM-1) mRNA, partial cds.	2133	100
611	AE003859	Xylella fastidiosa 9a5c	hypothetical protein	108	39
612	AK002185	Homo sapiens	FLJ11323 fis, clone PLACE1010362, weakly similar to 1-PHOSPHATIDYLINOSITOL PHOSPHODIESTERASE PRECURSOR (EC 3.1.4.10).	451	33
614	AAB41980	Homo sapiens	08-FEB-2001 31-MAR-2000 Human ORFX ORF1744 polypeptide sequence SEQ ID	116	76

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
.,,,,,	1,0		NO:3488.	 	Identity
615	AF161345	Homo sapiens	mRNA, partial cds.	439	100
616	AF116694	Homo sapiens	PRO2219	351	88
617	AAE03643	Homo sapiens	06-AUG-2001 05-DEC-2000 Human	1974	98
		•	extracellular matrix and cell adhesion		
			molecule-7 (XMAD-7).		
620	AL133640	Homo sapiens	cDNA DKFZp586C1021 (from clone DKFZp586C1021); partial cds.	2149	100
621	BC003369	Homo sapiens	ribosomal protein, large, P1, clone MGC:5215 IMAGE:2900846, mRNA, complete cds.	161	76
622	BC012124	Homo sapiens	clone MGC:20188 IMAGE:4564707, mRNA, complete cds.	810	100
625	AK008513	Mus musculus	putative	440	50
626	M32639	Homo sapiens	salivary statherin gene, exons 2-6.	276	87
627	BC008282	Homo sapiens	Similar to SH3-domain binding protein 1, clone MGC:10501 IMAGE:3639782, mRNA, complete cds.	897	96
628	AAG04000	Homo sapiens	06-OCT-2000 21-FEB-2000 Human secreted protein, SEQ ID NO: 8081.	515	100
629	AC011473	Homo sapiens	19, BAC BC349142 (CTC-518B2), complete sequence.	1392	100
632	AAY82615	Homo sapiens	02-AUG-2000 12-OCT-1998 Human PTHrP monoclonal antibody clone 1C1-3 protein SEQ ID NO:14.	768	88
633	AAB15539	Homo sapiens	28-FEB-2001 04-APR-2000 Human immune system molecule from Incyte clone 2907049.	637	98
634	AC018513	Homo sapiens	14 clone RP11-58H3 map 14q31, complete sequence.	818	100
635	X03249	Bos taurus	epsilon-4 beta-globin	321	79
636	AB046099	Macaca fascicularis	unnamed protein product	395	88
637	AC006033	Homo sapiens	clone RP11-121A8 from 7p14-p13, complete sequence.	1017	95
638	BC009488	Homo sapiens	Similar to CG10958 gene product, clone MGC:16372 IMAGE:3929220, mRNA, complete cds.	848	99
639	AL359620	Homo sapiens	cDNA DKFZp762P2111 (from clone DKFZp762P2111).	615	100
640	AB003184	Homo sapiens	for ISLR, complete cds.	880	59
641	AB036921	Pagrus major	maturation-inducing protein	797	69
643	AF284422	Homo sapiens	cotransporter-interacting protein mRNA, complete cds.	4694	100
646	AE000659	Homo sapiens	receptor alpha delta locus from bases 250472 to 501670 (section 2 of 5) of the Complete Nucleotide Sequence.	577	100
648	AAR59748	Homo sapiens	13-FEB-1995 14-DEC-1992 T cell receptor Valpha2.3 chain.	636	100
649	AJ004871	Homo sapiens	for TCR alpha chain, specific for Mage 3/HLA-A2.	1328	94
650	AF043179	Homo sapiens	cell receptor beta chain (TCRBV13S1- TCRBJ2S1) mRNA, complete cds.	1286	92
651	AAG74462	Homo sapiens	03-SEP-2001 28-SEP-2000 Human colon cancer antigen protein SEQ ID NO:5226.	143	75
652	AAE02653	Homo sapiens	06-AUG-2001 03-NOV-2000 Human gene 1 encoded uteroglobin-like protein from cDNA	287	98

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
			clone HTELR92.		
654	AAY70457	Homo sapiens	21-JUN-2000 02-SEP-1999 Human membrane channel protein-7 (MECHP-7).	1425	97
655	AJ406931	Homo sapiens	for keratin associated protein 3.1 (KRTAP3.1 gene).	598	100
656	AK000366	Homo sapiens	FLJ20359 fis, clone HEP16626.	2151	100
657	AF116688	Homo sapiens	PRO2133	370	98
658	BC002505	Homo sapiens	small nuclear ribonucleoprotein polypeptide F, clone MGC:1615 IMAGE:3051263, mRNA, complete cds.	222	84
659	D87009	Homo sapiens	lambda gene locus DNA, clone:288A10.	1822	99
660	AK000349	Homo sapiens	FLJ20342 fis, clone HEP13572.	3028	99
661	AK010756	Mus musculus	putative	653	84
662	AE006360	Lactococcus lactis subsp. lactis	HYPOTHETICAL PROTEIN	287	34
663	AC004832	Homo sapiens	clone RP4-539M6 from 22, complete sequence.	220	100
664	AB037902	Homo sapiens	AKR mRNA for truncated aldo-keto reductase type A, complete cds.	670	100
665	AF060511	Homo sapiens	016b10 My016 protein mRNA, complete cds.	133	52
666	M33014	Drosophila melanogaster	ubiquitin	153	62
667	AK022128	Homo sapiens	FLJ12066 fis, clone HEMBB1002266, moderately similar to NEURONAL PROTEIN.	1397	100
669	AL137512	Homo sapiens	cDNA DKFZp564E0178 (from clone DKFZp564E0178); partial cds.	751	100
670	S68015	human, mRNA, 1020 nt]. [Homo sapiens		1664	100
671	U89336	Homo sapiens	class III region containing NOTCH4 gene, partial sequence, homeobox PBX2 (HPBX) gene, receptor for advanced glycosylation end products (RAGE) gene, complete cds, and 6 unidentified cds, complete sequence.	2133	100
672	U89336	Homo sapiens	class III region containing NOTCH4 gene, partial sequence, homeobox PBX2 (HPBX) gene, receptor for advanced glycosylation end products (RAGE) gene, complete cds, and 6 unidentified cds, complete sequence.	2094	96
673	AL136746	Homo sapiens	cDNA DKFZp434K0512 (from clone DKFZp434K0512); complete cds.	962	94
674	AF125535	Homo sapiens	homolog mRNA, complete cds.	502	95
675	AF227130	Homo sapiens	taste receptor T2R3 gene, complete cds.	1629	100
677	AB046626	Macaca fascicularis	hypothetical protein	291	93
678	AC002077	Homo sapiens	cosmid clone LUCA17 from 3p21.3, complete sequence.	1145	100
679	AE000659	Homo sapiens	receptor alpha delta locus from bases 250472 to 501670 (section 2 of 5) of the Complete Nucleotide Sequence.	565	100
680	AAY99368	Homo sapiens	08-AUG-2000 01-SEP-1999 Human PRO1326 (UNQ686) amino acid sequence SEQ ID NO:100.	2034	100

SEQ ID NO:	Accession No.	Species	Description	Score	% Identity
682	BC000555	Homo sapiens	ribosomal protein L37a, clone MGC:1638 IMAGE:3162085, mRNA, complete cds.	187	55

TABLE 3

SEQ ID NO:	Accession No.	Description	Results*
343	BL00895	3-hydroxyisobutyrate	BL00895B 21.14 7.061e-22 151-190
		dehydrogenase proteins.	BL00895C 20.10 8.071e-22 200-236
			BL00895A 12.61 1.973e-18 42-63
351	PR00907	THROMBOMODULIN	PR00907B 11.29 9.299e-10 234-251
		SIGNATURE	
355	BL00585	Ribosomal protein S5 proteins.	BL00585A 28.43 1.391e-40 103-155
357	PR00078	GLYCERALDEHYDE-3-	PR00078B 7.45 3.250e-24 146-165
		PHOSPHATE	PR00078D 11.49 2.800e-21 232-250
		DEHYDROGENASE	PR00078E 10.50 6.211e-16 272-288
		SIGNATURE	PR00078C 15.99 8.000e-16 173-190
			PR00078A 10.38 1.000e-15 111-125
359	BL01282	BIR repeat proteins.	BL01282B 30.49 1.000e-13 523-562
361	BL00970	Nuclear transition protein 2	BL00970C 14.80 9.773e-09 70-108
		proteins.	
362	DM00191	w SPAC8A4.04C	DM00191A 8.16 9.640e-09 12-25
		RESISTANCE SPAC8A4.05C	
		DAUNORUBICIN.	<u></u>
365	PR00500	POLYCYSTIC KIDNEY	PR00500B 7.74 3.558e-09 396-417
		DISEASE PROTEIN	
		SIGNATURE	
367	BL50002	Src homology 3 (SH3) domain	BL50002B 15.18 1.600e-10 141-155
	ļ.,	proteins profile.	BL50002B 15.18 6.000e-09 42-56
368	BL50002	Src homology 3 (SH3) domain	BL50002B 15.18 1.600e-10 141-155
		proteins profile.	BL50002B 15.18 6.000e-09 42-56
369	BL00240	Receptor tyrosine kinase class	BL00240F 17.74 4.196e-11 552-600
		III proteins.	
370	BL01238	GDA1/CD39 family of	BL01238C 14.36 2.080e-16 212-234
		nucleoside phosphatases	BL01238D 10.19 1.180e-12 255-269
		proteins.	BL01238A 11.72 5.673e-11 86-101
371	PR00679	PROHIBITIN SIGNATURE	PR00679F 8.03 7.848e-25 122-146
			PR00679E 12.82 6.674e-18 97-117
			PR00679D 11.91 3.739e-16 74-91
	1		PR00679B 13.63 8.071e-16 28-48
	1		PR00679C 14.44 7.465e-14 51-70
	į		PR00679G 6.13 1.340e-13 157-174
374	DD 00700	DD OTED LEVE OGD IE	PR00679A 14.03 1.295e-12 10-27
3/4	PR00700	PROTEIN TYROSINE	PR00700D 12.47 4.462e-11 253-272
		PHOSPHATASE SIGNATURE	
375	PD00066	PROTEIN ZINC-FINGER	PD00066 13.92 2.385e-15 254-267
373	1 200000	METAL-BINDI.	PD00066 13.92 2.800e-14 310-323
		METAL-BINDI.	PD00066 13.92 7.429e-12 282-295
377	PR00925	NONHISTONE	PR00925B 3.73 6.625e-10 12-25
377	11(00)23	CHROMOSOMAL PROTEIN	FR00923B 3.73 6.023e-10 12-23
		HMG17 FAMILY	
		SIGNATURE	
378	PR00049	WILM'S TUMOUR PROTEIN	PR00049D 0.00 8.071e-10 3-18
5,0	, ,	SIGNATURE	1 100004712 0.00 0.0716-10 3-18
380	PF00084	Sushi domain proteins (SCR	PF00084B 9.45 3.250e-10 116-128
550	1100007	repeat proteins.	11 000040 7.43 3.2300-10 110-128
383	BL00636	Nt-dnaJ domain proteins.	BL00636A 8.07 1.947e-17 18-35
505	5200030	14-dias domain proteins.	BL00636B 15.11 5.500e-16 46-67
384	BL00636	Nt-dnaJ domain proteins.	BL00636A 8.07 1.947e-17 18-35
55.	2200030	11. dias domain proteins.	BL00636A 8.07 1.9476-17 18-33 BL00636B 15.11 5.500e-16 46-67
387	BL00741	Guanine-nucleotide	BL00741B 14.27 1.333e-14 302-325
	1 2000141	Guannic-naciconac	

SEQ ID NO:	Accession No.	Description	Results*
		dissociation stimulators CDC24 family sign.	
388	PF00628	PHD-finger.	PF00628 15.84 9.419e-09 179-194
392	PR00215	NEUROMODULIN	PR00215C 13.98 4.364e-09 201-222
		SIGNATURE	
394	PD00078	REPEAT PROTEIN ANK NUCLEAR ANKYR.	PD00078B 13.14 2.350e-10 132-145
397	BL01262	Eukaryotic initiation factor 1A proteins.	BL01262 22.38 6.625e-12 25-80
402	BL00056	Ribosomal protein S17 proteins.	BL00056A 28.90 3.769e-32 116-156 BL00056B 20.86 6.727e-23 164-188
403	BL00019	Actinin-type actin-binding domain proteins.	BL00019D 15.33 9.705e-13 296-326
409	PR00259	TRANSMEMBRANE FOUR FAMILY SIGNATURE	PR00259C 16.40 2.459e-21 78-107 PR00259A 9.27 2.846e-18 11-35 PR00259B 14.81 2.250e-17 51-78 PR00259D 13.50 2.756e-15 221-248
412	PD00066	PROTEIN ZINC-FINGER METAL-BINDI.	PD00066 13.92 2.385e-15 105-118 PD00066 13.92 4.462e-15 161-174 PD00066 13.92 1.600e-14 189-202 PD00066 13.92 1.500e-13 133-146 PD00066 13.92 1.500e-13 217-230 PD00066 13.92 1.000e-11 21-34 PD00066 13.92 2.957e-11 77-90
413	BL00028	Zinc finger, C2H2 type, domain proteins.	BL00028 16.07 3.400e-10 214-231 BL00028 16.07 7.171e-09 347-364
417	PF00791	Domain present in ZO-1 and Unc5-like netrin receptors.	PF00791B 28.49 8.057e-14 199-254 PF00791B 28.49 4.909e-11 166-221
421	BL00475	Ribosomal protein L15 proteins.	BL00475D 16.25 3.250e-19 130-152 BL00475C 13.06 3.700e-17 110-127 BL00475B 8.20 2.957e-11 36-46 BL00475A 10.62 8.560e-11 16-31
428	DM00215	PROLINE-RICH PROTEIN 3.	DM00215 19.43 2.286e-10 179-212
429	BL01153	NOL1/NOP2/sun family proteins.	BL01153D 19.69 4.375e-17 255-281 BL01153C 13.67 1.726e-11 205-219 BL01153A 13.77 4.300e-11 135-150
431	DM00984	w MYOD MYOBLAST DETERMINATION SHORT.	DM00984B 15.18 6.764e-17 142-197
441	PR00320	G-PROTEIN BETA WD-40 REPEAT SIGNATURE	PR00320C 13.01 2.800e-09 284-299 PR00320B 12.19 1.000e-08 146-161
443	PR00153	CYCLOPHILIN PEPTIDYL- PROLYL CIS-TRANS ISOMERASE SIGNATURE	PR00153A 12.98 1.667e-14 49-65 PR00153B 11.57 6.667e-12 78-91
444	PD02811	PROTEIN PEPTIDE REDUCTASE MG448 PILB FIMBRIA TRAN.	PD02811A 20.67 7.429e-12 4-42
446	PR00935	BAND 4.1 PROTEIN FAMILY SIGNATURE	PR00935D 10.20 4.656e-14 179-196 PR00935A 10.16 2.333e-12 40-53 PR00935C 11.98 2.500e-12 118-139 PR00935B 10.58 8.714e-11 105-119
447	BL00030	Eukaryotic RNA-binding region RNP-1 proteins.	BL00030A 14.39 1.643e-13 81-100
448	PR00401	SH2 DOMAIN SIGNATURE	PR00401B 12.94 7.333e-09 115-126 PR00401D 11.55 8.579e-09 144-155
453	BL00579	Ribosomal protein L29 proteins.	BL00579B 21.99 5.065e-21 35-65
457	BL00107	Protein kinases ATP-binding	BL00107A 18.39 4.960e-13 148-179

SEQ ID NO:	Accession No.	Description	Results*
		region proteins.	BL00107B 13.31 5.154e-12 222-238
458	BL00657	Fork head domain proteins.	BL00657A 19.39 1.191e-22 101-143
461	PF00615	Regulator of G protein	PF00615B 16.25 3.323e-14 103-120
463	BL00983	signalling domain proteins. Ly-6 / u-PAR domain proteins.	PF00615C 10.06 4.800e-10 180-194 BL00983C 12.69 6.885e-09 156-172
466	PR00358	BOMBESIN RECEPTOR	PR00358F 6.58 5.200e-09 15-29
100	1 K00550	SIGNATURE	1 FR00338F 0.38 3.2008-09 13-29
467	PD02784	PROTEIN NUCLEAR	PD02784B 26.46 1.000e-40 45-88
		RIBONUCLEOPROTEIN.	PD02784A 21.09 7.750e-37 5-42
			PD02784C 20.76 4.106e-09 97-143
469	BL00615	C-type lectin domain proteins.	BL00615A 16.68 2.080e-11 148-166
470	BL00615	C-type lectin domain proteins.	BL00615A 16.68 2.080e-11 175-193
475	PD01652	RECEPTOR CELL NK	PD01652B 8.50 7.207e-27 127-179
-	}	GLYCOPROTEIN	PD01652A 15.35 3.557e-17 137-173
400	PROCES	IMMUNOGLOB.	PD01652B 8.50 6.910e-10 32-84
478	PF00791	Domain present in ZO-1 and Unc5-like netrin receptors.	PF00791B 28.49 3.179e-12 40-95
479	PF00624	Flocculin repeat proteins.	DE00(241 0 10 7 165 00 071 201
480	PR00603	CYTOCHROME C1	PF00624I 9.10 7.165e-09 271-301
400	1 K00003	SIGNATURE	PR00603H 13.20 9.534e-09 285-301
482	BL01088	CAP protein.	BL01088F 14.83 5.404e-10 60-106
485	BL00412	Neuromodulin (GAP-43)	BL00412D 16.54 2.023e-11 45-96
		proteins.	BL00412D 16.54 3.204e-09 41-92
			BL00412D 16.54 5.684e-09 66-117
489	BL00353	HMG1/2 proteins.	BL00353A 9.60 1.000e-40 2-51
			BL00353B 11.47 1.000e-40 78-128
		1	BL00353C 14.83 1.000e-40 128-175
			BL00353A 9.60 5.661e-11 3-52
495	PF00523	Fusion glycoprotein F0.	PF00523D 11.39 7.188e-10 80-94
502	DM00031	IMMUNOGLOBULIN V REGION.	DM00031B 15.41 8.606e-11 78-112
505	PR00683	SPECTRIN PLECKSTRIN	PR00683D 15.87 9.864e-09 226-245
		HOMOLOGY DOMAIN SIGNATURE	
507	BL01189	Ribosomal protein S12e	BL01189A 14.27 7.513e-17 38-74
50.	DESTITO	proteins.	BL01189A 14.27 7.313e-17 38-74 BL01189A 14.27 5.245e-09 35-71
508	PD01094	ACID FATTY	PD01094D 7.35 7.094e-11 227-281
	1301071	DESATURASE	1 0010340 7.33 7.0346-11 227-281
		ENDOPLASMI.	
512	BL00028	Zinc finger, C2H2 type,	BL00028 16.07 2.286e-09 353-370
_		domain proteins.	
513	BL00028	Zinc finger, C2H2 type,	BL00028 16.07 2.286e-09 353-370
		domain proteins.	
514	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 5.714e-16 117-148
516	BL00951	ER lumen protein retaining	BL00951C 19.35 1.000e-40 93-142
		receptor proteins.	BL00951B 14.23 4.300e-31 38-69
			BL00951D 13.94 1.783e-30 142-177
			BL00951A 15.10 1.818e-29 2-38
517	BL00951	ER lumen protein retaining	BL00951D 13.94 2.761e-30 89-124
		receptor proteins.	BL00951A 15.10 1.818e-29 2-38
			BL00951B 14.23 5.950e-27 38-69
			BL00951C 19.35 4.493e-22 40-89
522	PF01105	emp24/gp25L/p24 family.	PF01105B 25.12 3.928e-12 176-228
526	BL00518	Zinc finger, C3HC4 type	BL00518 12.23 2.714e-10 31-40
524	DD00202	(RING finger), proteins.	DD000000 10 06 1 05
534	PD00787	SYNTHASE BIOSYNTHESIS	PD00787B 13.26 1.574e-09 91-105

SEQ ID NO:	Accession No.	Description	Results*
		TRANSFERASE.	
538	PF00632	HECT-domain (ubiquitin-	PF00632C 20.66 1.540e-20 554-586
		transferase).	PF00632B 18.45 8.313e-20 499-527
541	BL00478	LIM domain proteins.	BL00478B 14.79 9.679e-13 62-77
	:		BL00478B 14.79 5.750e-12 182-197
			BL00478B 14.79 6.500e-12 245-260
			BL00478B 14.79 3.400e-11 123-138
543	DM00547	1 kw CHROMO	DM00547F 23.43 6.538e-36 628-675
		BROMODOMAIN SHADOW	DM00547E 13.94 2.400e-18 387-410
	İ	GLOBAL.	DM00547C 17.30 9.486e-16 266-288
			DM00547B 11.28 9.217e-15 237-251
	İ		DM00547D 11.60 4.951e-12 357-371
545	PF00777	Sialyltransferase family.	DM00547A 12.38 6.455e-11 216-228
550	PD00066	PROTEIN ZINC-FINGER	PF00777C 18.60 5.291e-21 78-133 PD00066 13.92 3.769e-15 459-472
330	FD00000	METAL-BINDI.	PD00066 13.92 3.7696-13 439-472 PD00066 13.92 2.8006-14 206-219
		WETAL-BINDI.	PD00066 13.92 2.800e-14 206-219
	1	_	PD00066 13.92 2.800e-14 234-247
			PD00066 13.92 2.800e-14 431-444
		1	PD00066 13.92 2.800e-14 487-500
	•		PD00066 13.92 3.400e-14 375-388
			PD00066 13.92 5.200e-14 319-332
	Į.		PD00066 13.92 8.800e-14 403-416
			PD00066 13.92 4.000e-13 150-163
	-)(-		PD00066 13.92 5.500e-13 515-528
			PD00066 13.92 7.652e-11 262-275
553	PF00615	Regulator of G protein	PF00615B 16.25 8.839e-14 101-118
		signalling domain proteins.	PF00615C 10.06 3.700e-13 178-192
555	PR00180	CELLULAR	PR00180A 10.11 1.875e-16 75-98
}		RETINALDEHYDE-	PR00180D 12.78 1.155e-15 233-253
	İ	BINDING PROTEIN	PR00180B 16.42 4.493e-13 124-149
557	BL00018	SIGNATURE	PR00180C 10.92 2.901e-12 200-222
337	BL00018	EF-hand calcium-binding domain proteins.	BL00018 7.41 4.150e-10 494-507
559	BL01172	Ribosomal protein L44e	BL01172B 14.10 1.000e-40 15-57
		proteins.	BL01172C 16.78 3.400e-33 63-102
			BL01172A 7.78 3.520e-13 2-13
562	DM00031	IMMUNOGLOBULIN V	DM00031B 15.41 1.000e-10 83-117
		REGION.	
563	BL00484	Thyroglobulin type-1 repeat	BL00484B 9.04 6.344e-14 103-117
		proteins proteins.	BL00484C 17.01 8.125e-14 123-138
565	PF00566	Probable rabGAP domain	PF00566A 12.64 9.667e-10 111-121
		proteins.	PF00566B 11.92 1.300e-09 153-159
566	BL00580	Ribosomal protein L32e	BL00580A 17.63 9.899e-09 14-50
569	BL00674	proteins. AAA-protein family proteins.	PI 00674D 22 41 4 626 15 622 646
1 309	DL000/4	AAA-protein family proteins.	BL00674D 23.41 4.696e-15 599-646 BL00674B 4.46 1.333e-14 508-530
			BL00674B 4.46 1.333e-14 508-530 BL00674C 22.60 3.786e-14 541-584
572	BL00397	Site-specific recombinases	BL00397D 19.54 8.163e-10 279-299
L		proteins.	0.1036-10 2/9-299
575	BL00242	Integrins alpha chain proteins.	BL00242E 9.03 1.375e-26 1143-1172
			BL00242C 16.86 2.324e-23 483-513
			BL00242D 13.57 5.200e-22 570-595
			BL00242B 8.13 6.478e-11 394-404
			BL00242A 13.80 7.000e-11 75-87
500	DI 00415		BL00242D 13.57 3.957e-10 632-657
582	BL00415	Synapsins proteins.	BL00415N 4.29 2.445e-09 386-430

SEQ ID NO:	Accession No.	Description	Results*
583	PD00066	PROTEIN ZINC-FINGER	PD00066 13.92 1.000e-14 165-178
		METAL-BINDI.	PD00066 13.92 5.800e-14 193-206
			PD00066 13.92 9.000e-13 221-234
			PD00066 13.92 1.000e-12 137-150
			PD00066 13.92 5.286e-12 249-262
			PD00066 13.92 9.143e-12 109-122
			PD00066 13.92 2.957e-11 81-94
585	BL50058	G-protein gamma subunit	BL50058 27.23 8.393e-31 35-83
		profile.	
587	PF00628	PHD-finger.	PF00628 15.84 6.806e-09 77-92
591	PR00450	RECOVERIN FAMILY	PR00450C 12.22 5.364e-12 65-87
		SIGNATURE	
592	PR00450	RECOVERIN FAMILY	PR00450C 12.22 5.364e-12 65-87
		SIGNATURE	
600	BL00617	RecF protein.	BL00617A 25.53 6.308e-11 61-104
603	PR00216	OSTEOPONTIN	PR00216C 9.63 8.636e-09 189-215
		SIGNATURE	
604	BL00019	Actinin-type actin-binding	BL00019D 15.33 7.660e-17 397-427
		domain proteins.	
610	PF00855	PWWP domain proteins.	PF00855 13.75 7.000e-10 414-431
613	BL01228	Hypothetical cof family	BL01228D 17.44 2.523e-10 609-634
		proteins.	
629	BL00021	Kringle domain proteins.	BL00021B 13.33 4.240e-16 48-66
635	BL01033	Globins profile.	BL01033B 13.81 5.500e-14 38-50
638	PF00992	Troponin.	PF00992A 16.67 7.868e-09 7-42
639	PD00066	PROTEIN ZINC-FINGER	PD00066 13.92 8.800e-14 50-63
		METAL-BINDI.	120000 1332 3.3300 1130 33
640	PR00500	POLYCYSTIC KIDNEY	PR00500B 7.74 7.964e-12 182-203
		DISEASE PROTEIN	
		SIGNATURE	
641	PD00066	PROTEIN ZINC-FINGER	PD00066 13.92 6.143e-12 316-329
		METAL-BINDI.	PD00066 13.92 6.192e-10 344-357
643	PD01941	TRANSMEMBRANE	PD01941A 14.81 2.662e-34 82-136
		COTRANSPORTER SYMP.	PD01941B 15.02 2.246e-28 267-314
		1	PD01941D 27.18 9.194e-19 501-550
			PD01941C 19.96 6.786e-13 347-402
649	DM00031	IMMUNOGLOBULIN V	DM00031B 15.41 3.278e-09 79-113
		REGION.	
650	BL00290	Immunoglobulins and major	BL00290A 20.89 8.200e-12 162-185
		histocompatibility complex	
		proteins.	
654	BL00407	Connexins proteins.	BL00407E 22.17 1.000e-40 164-209
			BL00407B 14.23 7.231e-35 39-70
			BL00407A 18.57 5.250e-29 2-39
		ĺ	BL00407C 14.61 7.097e-28 70-98
			BL00407D 17.61 4.000e-25 125-155
656	PR00359	B-CLASS P450 SIGNATURE	PR00359F 24.20 4.536e-10 310-338
661	BL01064	Pyridoxamine 5'-phosphate	BL01064C 15.22 1.205e-09 307-340
		oxidase proteins.	
664	PR00069	ALDO-KETO REDUCTASE	PR00069A 16.01 1.000e-18 42-67
		SIGNATURE	PR00069B 11.33 1.735e-13 102-121
665	PD02462	PROTEIN BOLA	PD02462A 22.48 9.873e-12 13-48
		TRANSCRIPTION	
		REGULATION AC.	
666 ,	PR00348	UBIQUITIN SIGNATURE	PR00348A 7.86 8.625e-09 11-32
667	BL01052	Calponin family repeat	BL01052B 15.31 2.518e-10 511-537
		proteins.	1

SEQ ID NO:	Accession No.	Description	Results*
671	PD02327	GLYCOPROTEIN ANTIGEN	PD02327B 19.84 8.941e-23 143-165
		PRECURSOR	PD02327A 8.89 1.000e-13 115-127
		IMMUNOGLO.	PD02327C 15.47 5.500e-13 209-224
672	PD02327	GLYCOPROTEIN ANTIGEN	PD02327B 19.84 8.941e-23 159-181
		PRECURSOR	PD02327A 8.89 1.000e-13 115-127
		IMMUNOGLO.	PD02327C 15.47 5.500e-13 225-240
678	PR00441	G-PROTEIN ALPHA	PR00441B 16.16 4.667e-26 163-186
		SUBUNIT GROUP I	PR00441C 14.17 1.409e-24 192-210
		SIGNATURE	PR00441A 10.69 1.375e-19 31-47

^{*} Results include in order: Accession No., subtype, e-value, and amino acid position of the signature in the corresponding polypeptide

TABLE 4

SEQ ID NO:	Pfam Model	Description	E-value	Score
350	K tetra	K+ channel tetramerisation domain	2.3e-31	117.6
351	zona_pellucida	Zona pellucida-like domain	2.2e-25	97.7
355	Ribosomal S5	Ribosomal protein S5	1.7e-46	167.9
357	gpdh	Glyceraldehyde 3-phosphate dehydrogenase, NA	3.le-144	349.8
429	Nol1 Nop2 Sun	NOL1/NOP2/sun family	4.5e-19	68.6
431	LIM	LIM domain	8.6e-32	119.1
441	WD40	WD domain, G-beta repeat	2.3e-07	37.9
443	pro_isomerase	Cyclophilin type peptidyl-prolyl cis-tr	5.3e-34	120.4
444	DUF25	Domain of unknown function DUF25	1.1e-11	46.9
446	Band_41	FERM domain (Band 4.1 family)	3.2e-77	242.4
447	rrm	RNA recognition motif.	1.1e-33	125.4
448	SH2	SH2 domain	1.7e-33	100.2
449	UIM	Ubiquitin interaction motif	0.00071	26.3
453	Ribosomal L29	Ribosomal L29 protein	1.7e-15	64.9
454	NTF2	Nuclear transport factor 2 (NTF2) domain	3.2e-07	37.4
457	pkinase	Protein kinase domain	6e-40	146.1
458	Fork head	Fork head domain	1e-28	108.8
460	PC4	Transcriptional Coactivator p15 (PC4)	2.1e-38	141.0
461	RGS	Regulator of G protein signaling domain	2.6e-45	164.0
465	COX7a	Cytochrome c oxidase subunit VIIa	2.3e-40	147.5
467	rrm	RNA recognition motif.	3.2e-15	64.0
469	lectin c	Lectin C-type domain	5.1e-06	33.3
470	lectin c	Lectin C-type domain	5.1e-06	33.3
475	ig	Immunoglobulin domain	9.1e-07	26.9
478	ank	Ank repeat	3e-15	64.1
481	Zip	ZIP Zinc transporter	3.8e-31	116.9
489	HMG_box	HMG (high mobility group) box	8e-53	188.9
490	PH	PH domain	2.8e-13	52.3
494	Ulp1 C	Ulp1 protease family, C-terminal catalytic d	1.2e-11	52.1
495	Peptidase_C6	Helper component proteinase	0.0056	7.9
502	ig	Immunoglobulin domain	2.3e-09	35.2
503	ig	Immunoglobulin domain	9.2e-09	33.3
505	PH	PH domain	1.9e-14	56.4
507	Ribosomal_L7Ae	Ribosomal protein L7Ae/L30e/S12e/Gadd4	8.2e-14	59.3
512	zf-C2H2	Zinc finger, C2H2 type	1.1e-10	48.9
513	zf-C2H2	Zinc finger, C2H2 type	3.2e-16	67.3
514	pkinase	Protein kinase domain	3.4e-26	98.4
516	ER_lumen_recept	ER lumen protein retaining receptor	3.5e-144	492.4
517	ER_lumen_recept	ER lumen protein retaining receptor	1.8e-88	307.3
522	EMP24_GP25L	emp24/gp25L/p24 family	6.9e-06	28.1
526	SPRY	SPRY domain	2.3e-30	114.3
538	HECT	HECT-domain (ubiquitin-transferase)	1.1e-115	397.8
540	Rhomboid	Rhomboid family	4.2e-42	153.3
541	LIM	LIM domain	2e-35	131.1
542	Glycos_transf_2	Glycosyl transferase	1.7e-25	98.1
543	SNF2_N	SNF2 and others N-terminal domain	5.9e-104	358.8
545	Glyco_transf_29	Glycosyltransferase family 29	7.3e-20	79.4
546	LysM	LysM domain	5e-06	33.5
550	zf-C2H2	Zinc finger, C2H2 type	1.1e-104	361.2
553	RGS	Regulator of G protein signaling domain	5.1e-52	186.2
554	TBC	TBC domain	7.2e-35	129.3
555	CRAL_TRIO	CRAL/TRIO domain	4.5e-47	158.6
559	Ribosomal_L44	Ribosomal protein L44	1e-48	175.3
561	TIR	TIR domain	0.063	9.9

SEQ ID NO:	Pfam Model	Description	E-value	Score
562	ig	Immunoglobulin domain	3.5e-08	31.4
563 565	thyroglobulin_1	Thyroglobulin type-1 repeat	3.9e-24	93.6
	TBC	TBC domain	1.2e-54	195.0
568	zf-C2H2	Zinc finger, C2H2 type	7.1e-08	39.6
569	AAA	ATPase family associated with various cellul	2e-44	161.0

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PDB annotation		OXIDOREDUCTASE (D, L) STERFOSPECIFIC OPINE	DEHYDROGENASE.	OXIDOREDUCTASE		OXIDOREDUCTASE AMINO ACID	DEHYDROGENASE, OXIDATIVE	DEAMINATION MECHANISM, 2	OXIDOREDUCTASE			OXIDOREDUCTASE	OXYDOREDUCTASE			OXIDOREDUCTASE ROSSMANN	FOLD, TERNARY COMPLEX,	CRYSTALLOGRAPHIC DIMER	OXIDOREDUCTASE	DEHYDROGENASE, ALCOHOL,	NICOTINAMIDE COENZYME,	STEROID 2 BINDING	OXIDOREDUCTASE HCDH; ABORTIVE	TERNARY COMPLEX		OXIDOREDUCTASE SHORT-CHAIN	DEHYDROGENASE/REDUCTASE, BILE	ACID CATABOLISM			
Coumpound		N-(1-D- CARBOXYI ETHYI)-I -	NORVALINE	DEHYDROGENASE;	CHAIN: NULL;	L-PHENYLALANINE	DEHYDROGENASE;	CHAIN: A; L-	PHENYLALANINE	DEHYDROGENASE;	CHAIN: B;	GLYCERALDEHYDE-3-	PHOSPHATE	DEHYDROGENASE;	CHAIN: P, R, O, Q;	UDP-GLUCOSE	DEHYDROGENASE;	CHAIN: A;	ALCOHOL	DEHYDROGENASE;	CHAIN: A, B;		L-3-HYDROXYACYL-COA	DEHYDROGENASE;	CHAIN: A, B;	7 ALPHA-	HYDROXYSTEROID	DEHYDROGENASE;	CHAIN: A, B;	OXIDOREDUCTASE(CHOH	(D)-NAD(P)+(A)) D-
SeqFold	3										İ																				_
PMF	200	1.00				0.31						0.82				-0.01			0.39				0.10			0.31				-0.03	
Verify Score		0.52				0.58						0.54				0.14			0.10				0.31			0.32				0.17	
PSI RY AST	Score	3.4e-11				1.7e-10						1.4e-06				1e-36			8.5e-06			- 1	1.2e-43			8.5e-06				1.7e-29	
End		158				198						112				333			91			-†	324			135				205	
Start		40				37						40				40			39				40			37				25	
Chain ID	}					4						Д				٧	_		A				4							∢	
PDB ID		1bg6				lcld						Ictz		_		iệi —			lee2				16y			1 fmc				lgdh	_
SEQ ID	NO:	343				343						343				343			343				343			343				343	

PDB annotation			OXIDOREDUCTASE		OXIDOREDUCTASE 6PGDH, 6-PGDH; OXIDOREDUCTASE, CHOH(D)- NADP+(B)	OXIDOREDUCTASE, OXIDOREDUCTASE, OXIDOREDUCTASE, NAD	
Coumpound	GLYCERATE DEHYDROGENASE (APO FORM) (E.C.1.1.1.29) 1GDH 3	OXIDOREDUCTASE(CHOH (D)-NAD(A)) APO-*L- *LACTATE DEHYDROGENASE (E.C.1.1.27) 1LDB 4	LEUCINE DEHYDROGENASE; CHAIN: A, B;	OXIDOREDUCTASE(CHOH (D)-NAD (A)) L-LACTATE DEHYDROGENASE (E.C.1.1.27) (T-STATE) MUTANT 1LLD 3 WITH CYS 199 REPLACED BY SER (C199S) COMPLEX WITH NADH 1LLD 4	6-PHOSPHOGLUCONATE DEHYDROGENASE; CHAIN: A, B;	L-ALANINE DEHYDROGENASE; CHAIN: A;	OXIDOREDUCTASE (NAD(A)) D-3- PHOSPHOGLYCERATE DEHYDROGENASE (PHOSPHOGLYCERATE IPSD 3 DEHYDROGENASE) (E.C.1.1.95) IPSD-4
SeqFold Score							
PMF Score	i	0.30	0.23	0.18	0.99	09.0	0.48
Verify Score		0.02	0.34	0.17	0.27	0.36	0.15
PSI BLAST Score		1.2e-05	1.7e-08	1.7e-06	3.4e-37	1e-09	5.1e-34
End		126	226	156	307	133	213
Start		39	23	41	40	41	18
Chain ID			А	A	4	A	Ą
PDB U		1ldb	11ch	11114	lpgi	1pjc	1 psd
SEQ ID NO:		343	343	343	343	343	343

SEQ ID NO:	PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
343	19p8	V V	37	212	le-21	0.09	-0.09		FORMATE DEHYDROGENASE; CHAIN: A, B;	OXIDOREDUCTASE SIMILAR TO THE PREVIOUSLY SOLVED FORMATE DEHYDROGENASE, 2 OXIDOREDUCTASE
343	2cmd		40	123	5.1e-06	0.01	0.07		OXIDOREDUCTASE(NAD(A)-CHOH(D)) MALATE DEHYDROGENASE (E.C.1.1.37) 2CMD 3	
343	2did	A	01	162	5.4e-18	0.23	0.59		D-LACTATE DEHYDROGENASE; 2DLD 5 CHAN: A, B; 2DLD 6	OXIDOREDUCTASE (CHOH(D)- NAD+(A)) R-LACTATE DEHYDROGENASE; 2DLD 7
343	2dld	٧	29	215	le-34	0.30	0.29		D-LACTATE DEHYDROGENASE; 2DLD 5 CHAIN: A, B; 2DLD 6	OXIDOREDUCTASE (CHOH(D)- NAD+(A)) R-LACTATE DEHYDROGENASE; 2DLD 7
343	2pgd		42	328	1.7e-45	0.13	0.89		OXIDOREDUCTASE (CHOH(D)-NADP+(A)) 6- PHOSPHOGLUCONATE DEHYDROGENASE (6- PGDH) (E.C.1.1.1.44) 2PGD 3	
343	3hdh	∢	36	329	8.5e-43			59.63	L-3-HYDROXYACYL COA DEHYDROGENASE; CHAIN: A, B, C;	OXIDOREDUCTASE SCHAD; OXIDOREDUCTASE, BETA OXIDATION, SCHAD, CATALYTIC ACTIVITY: 2 L-3-HYDROXYACYL- COA + NAD(+) = 3-OXOACYL-COA + NADH
343	3hdh	V	40	324	8.5e-43	0.15	0.23		L-3-HYDROXYACYL COA DEHYDROGENASE; CHAIN: A, B, C;	OXIDOREDUCTASE SCHAD; OXIDOREDUCTASE, BETA OXIDATION, SCHAD, CATALYTIC ACTIVITY: 2 L-3-HYDROXYACYL- COA + NAD(+) = 3-OXOACYL-COA + NADH
343	3hdh	ပ	15	278	6.8e-32			75.95	L-3-HYDROXYACYL COA DEHYDROGENASE;	OXIDOREDUCTASE SCHAD; OXIDOREDUCTASE, BETA

SEQ FIG	PDB ID	Chain ID	Start	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									CHAIN: A, B, C;	OXIDATION, SCHAD, CATALYTIC ACTIVITY: 2 L-3-HYDROXYACYL- COA + NAD(+) = 3-0XOACYL-COA + NADH
343	3hdh	U	40	317	6.8e-32	0.12	0.01		L-3-HYDROXYACYL COA DEHYDROGENASE; CHAIN: A, B, C;	OXIDOREDUCTASE SCHAD; OXIDOREDUCTASE, BETA OXIDATION, SCHAD, CATALYTIC ACTIVITY: 2 L-3-HYDROXYACYL- COA + NAD(+) = 3-OXOACYL-COA + NADH
345	1b3u	A	272	582	0.00016	0.12	0.39		PROTEIN PHOSPHATASE PP2A; CHAIN: A, B;	SCAFFOLD PROTEIN SCAFFOLD PROTEIN, PP2A, PHOSPHORYLATION, HEAT REPEAT
345	3bct		208	639	7.2e-14	-0.10	90.0		BETA-CATENIN; CHAIN: NULL;	ARMADILLO REPEAT ARMADILLO REPEAT, BETA-CATENIN, CYTOSKELETON
\neg										
346	lari	ပ	128	215	3.4e-16	90:00	-0.19		CYTOCHROME C OXIDASE; CHAIN: A, B; ANTIBODY FV FRAGMENT; CHAIN: C, D;	COMPLEX (OXIDOREDUCTASE/ANTIBODY) CYTOCHROME AA3, COMPLEX IV, FERROCYTOCHROME C, COMPLEX (OXIDOREDUCTASE/ANTIBODY), ELECTRON TRANSPORT, 2 TRANSMEMBRANE, CYTOCHROME OXIDASE, ANTIBODY COMPLEX
346	1dq!	н		218	3.4e-16	0.04	-0.20		IGM MEZ IMMUNOGLOBULIN; CHAIN: L; IGM MEZ IMMUNOGLOBULIN; CHAIN: H;	IMMUNE SYSTEM IMMUNOGLOBULIN FOLD, ANTIBODY, IGM, FV
	1dsf	н	130	214	1.7e-16	0.10	-0.17		ANTICANCER ANTIBODY BI; CHAIN: L, H;	IMMUNOGLOBULIN BIDSFV; MONOCLONAL ANTIBODY, ANTITUMOR, IMMUNOGLOBULIN

PDB annotation			IMMUNOGLOBULIN NMR, VH DOMAIN, ANTIBODY, HUMAN, IMMUNOGLOBULIN	STRUCTURAL PROTEIN INTEGRIN- BINDING PROTEIN INV GENE	 		STRUCTURAL PROTEIN INTEGRIN- BINDING PROTEIN INV GENE	╫
Coumpound	IMMUNOGLOBULIN IMMUNOGLOBULIN M (IG-M) FV FRAGMENT IIGM 3	IMMUNOGLOBULIN FV FRAGMENT (MURINE SE155-4) COMPLEX WITH THE TRISACCHARIDE: IMFA 3 ALPHA-D- GALACTOSE(1-2)[ALPHA- D-ABEQUOSE(1-3)]ALPHA- IMFA 4 D-MANNOSE (PI- OME) (PART OF THE CELL-SURFACE CARBOHYDRATE IMFA 5 OF PATHOGENIC SALMONELLA) IMFA 6	VH-P8; CHAIN: NULL;	INVASIN; CHAIN: A;	GLYCOSYLTRANSFERASE CYCLODEXTRIN GLUCANOTRANSFERASE (E.C.2.4.1.19) (CGTASE) ICYG 3	VIRUS TOMATO BUSHY STUNT VIRUS 2TBV 4	NVASN; CHAIN: A;	GLYCOSYLTRANSFERASE
SeqFold Score				79.67		59.57	19.61	
PMF Score	-0.19	6.18	-0.18		-0.09			-0.09
Verify Score	0.09	0.02	0.37		0.02			0.02
PSI BLAST	le-15	16-19	1.2e-16	1.8e-24	1.4e-15	1.8e-20	1.8e-24	1.4e-15
End	214	220	214	464	411	395	464	411
Start AA	128	44	128	9	57	74	9	57
Chain ID	H			A		ပ	A	
PDB CI	ligm	Imfa	lvhp	lcwv	lcyg	2tbv	Icwv	lcyg
SEQ El SE	346	346	346	347	347	347	348	348

PDB annotation			POTASSIUM CHANNELS POTASSIUM CHANNELS, TETRAMERIZATION DOMAIN, X-RAY 2 STRUCTURE, APLYSIA KV1.1	POTASSIUM CHANNELS POTASSIUM CHANNELS, TETRAMERIZATION DOMAIN, X-RAY 2 STRUCTURE, APLYSIA KV1.1	GENE REGULATION POZ DOMAIN; PROTEIN-PROTEIN INTERACTION DOMAIN, TRANSCRIPTIONAL 2 REPRESSOR, ZINC-FINGER PROTEIN, X-RAY CRYSTALLOGRAPHY, 3 PROTEIN STRUCTURE, PROMYELOCYTIC LEUKEMIA, GENE REGULATION	SIGNALING PROTEIN VOLTAGE- GATED POTASSIUM CHANNEL, ASSEMBLY DOMAIN, TETRAMER	METAL TRANSPORT ION CHANNEL, OXIDOREDUCTASE, BETA SUBUNIT	SIGNALING PROTEIN VOLTAGE- GATED POTASSIUM CHANNEL, TETRAMERIZATION DOMAIN, 2
Coumpound	CYCLODEXTRIN GLUCANOTRANSFERASE (E.C.2.4.1.19) (CGTASE) 1CYG 3	VIRUS TOMATO BUSHY STUNT VIRUS 2TBV 4	POTASSIUM CHANNEL KV1.1; CHAIN: NULL;	POTASSIUM CHANNEL KV1.1; CHAIN: NULL;	PROMYELOCYTIC LEUKEMIA ZINC FINGER PROTEIN PLZF; CHAIN: A;	KVI.2 VOLTAGE-GATED POTASSIUM CHANNEL; CHAIN: A, B, C, D, E, F, G, H;	KV BETA2 PROTEIN; CHAIN: A; POTASSIUM CHANNEL KV1.1; CHAIN: E;	KVI.2 VOLTAGE-GATED POTASSIUM CHANNEL; CHAIN: A, B, C, D;
SeqFold Score		59.57						
PMF Score			0.49	0.49	0.87	0.39	0.54	0.45
Verify Score			-0.21	-0.39	0.47	-0.31	0.02	-0.06
PSI BLAST Score		1.8e-20	1.7e-14	1.1e-24	5.1e-17	1.4e-14	le-14	le-14
End		395	124	123	136	124	124	124
Start AA		74	41	42	34	41	39	41
Chain ID	-	၁			⋖	¥	ம	٧
PDB ID		2tbv	1a68	1a68	1buo	ldsx	lexb	Iqdv
SEQ ID NO:		348	350	350	350	350	350	350

SEQ ID NO:	PDB ID	Chain ID	Start	End AA	· PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
[5]					3000					INTRACELLULAR GATE, TETRAMER
350	1qdv	¥.	42	132	1.1e-27	-0.18	0.42		KVI.2 VOLTAGE-GATED POTASSIUM CHANNEL; CHAIN: A, B, C, D;	SIGNALING PROTEIN VOLTAGE- GATED POTASSIUM CHANNEL, TETRAMERIZATION DOMAIN, 2 NTP A CEI I III A P GATE TETP AMED
350	ptit	4	41	124	1.7e-14	-0.09	0.58		POTASSIUM CHANNEL KV1.1; CHAIN: A;	PROTON TRANSPORT POTASSIUM CHANNELS, TETRAMERIZATION DOMAIN, X-RAY STRUCTURE, 2 APLYSIA KVIII. PROTON TRANSPORT
350	3kvt		40	132	5.4e-24	0.34	0.96		POTASSIUM CHANNEL PROTEIN SHAW; CHAIN: NULL;	POTASSIUM CHANNEL POTASSIUM CHANNEL, TETRAMERIZATION DOMAIN, MOLECULAR 2 RECOGNITION, ZINC-BINDING
350	3kvt		41	140	1e-16	0.37	0.98		POTASSIUM CHANNEL PROTEIN SHAW; CHAIN: NULL;	POTASSIUM CHANNEL POTASSIUM CHANNEL, TETRAMERIZATION DOMAIN, MOLECULAR 2 RECOGNITION, ZINC-BINDING
351	laut	,i	22.7	279	5.4e-20	0.08	0.40		ACTIVATED PROTEIN C; CHAIN: C, I.; D-PHE-PRO- MAI; CHAIN: P; COAGULATION FACTOR VIIA (LIGHT CHAIN) (DES- GLA); CHAIN: L; COAGULATION FACTOR VIIA (HEAVY CHAIN) (DES-GLA); CHAIN: H; DEGR-CK INHIBITOR (GLU-GLY-ARM); CHAIN:	COMPLEX (BLOOD COAGULATION/INHIBITOR) AUTOPROTHROMBIN IIA; HYDROLASE, SERINE PROTEINASE), PLASMA CALCIUM BINDING, 2 GLYCOPROTEIN, COMPLEX (BLOOD COAGULATION/INHIBITOR) HYDROLASE BLOOD COAGULATION, FACTOR VIIA, SERINE PROTEASE, EGF, 2 INHIBITOR, CRYSTAL STRUCTURE
			_						J.	

03	BOG	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
NO S	en .	a l	AA	AA	BLAST Score	Score	Score	Score		
351	ldan	1	124	234	5.4e-18	-0.17	0.01		BLOOD COAGULATION FACTOR VIIA; CHAIN: L, H; SOLUBLE TISSUE	BLOOD COAGULATION, SERINE PROTEASE, COMPLEX, CO-FACTOR, 2 RECEPTOR ENZYME, INHIBITOR.
									FACTOR; CHAIN: T, U; D-	GLA, EGF, 3 COMPLEX (SERINE
									PHE-PHE-ARG-	PROTEASE/COFACTOR/LIGAND)
									(DFFRCMK) WITH CHAIN: C:	
351	Idan	ı	159	268	1.8e-22	0.18	0.17		BLOOD COAGULATION FACTOR VIIA; CHAIN: L, H: SOLUBLE TISSUE	BLOOD COAGULATION, SERINE PROTEASE, COMPLEX, CO-FACTOR, 2 RECEPTOR ENZYME, INHIBITOR.
									FACTOR; CHAIN: T, U; D-PHE-PHE-ARG-	GLA, EGF, 3 COMPLEX (SERINE PROTEASE/COFACTOR/LIGAND)
									CHLOROMETHYLKETONE (DFFRCMK) WITH CHAIN: C:	
351	Idan	1	182	273	3.4e-15	0.57	0.87		BLOOD COAGULATION FACTOR VIA: CHAIN: 1	BLOOD COAGULATION, SERINE
						_			H; SOLUBLE TISSUE	RECEPTOR ENZYME, INHIBITOR,
									FACTOR; CHAIN: T, U; D-PHE-PHE-ARG-	GLA, EGF, 3 COMPLEX (SERINE PROTEASE/COFACTOR/LIGAND)
									CHLOROMETHYLKETONE	
		-							(DFFRCMK) WITH CHAIN: C;	
351	Idva	L	179	268	7.2e-19	-0.07	0.43		DES-GLA FACTOR VIIA	HYDROLASE/HYDROLASE INHIBITOR
									(HEAVY CHAIN); CHAIN: H, I; DES-GLA FACTOR	PROTEIN-PEPTIDE COMPLEX
	-								VIIA (LIGHT CHAIN);	
									CHAIN: L, M; (DPN)-PHE-	
									ARG; CHAIN: C, D;	
									PEPTIDE E-76; CHAIN: X, v.	
351	ldva	T	182	273	3.4e-15	0.64	0.59		DES-GLA FACTOR VIIA	HYDROLASE/HYDROLASE INHIBITOR
7									(HEAVY CHAIN); CHAIN:	PROTEIN-PEPTIDE COMPLEX

PDB annotation		SERINE PROTEINASE COAGULATION FACTOR II; COAGULATION FACTOR II; FETOMODULIN, TM, CD141 ANTIGEN; EGR-CMK SERINE PROTEINASE, EGF-LIKE DOMAINS, ANTICOAGULANT COMPLEX, 2 ANTIFIBRINOLYTIC COMPLEX	SERINE PROTEINASE COAGULATION FACTOR II; COAGULATION FACTOR II; FETOMODULIN, TM, CD141 ANTIGEN; EGR-CMK SERINE PROTEINASE, EGF-LIKE DOMAINS, ANTICOAGULANT COMPLEX, 2 ANTIFIBRINOLYTIC COMPLEX	SERINE PROTEINASE COAGULATION FACTOR II; COAGULATION FACTOR II; FETOMODULIN, TM, CD141 ANTIGEN; EGR-CMK SERINE PROTEINASE, EGF-LIKE DOMAINS, ANTICOAGULANT COMPLEX, 2 ANTIFIBRINOLYTIC COMPLEX	MATRIX PROTEIN EXTRACELLULAR
Coumpound	H, I; DES-GLA FACTOR VIIA (LIGHT CHAIN); CHAIN: L, M; (DPN)-PHE- ARG; CHAIN: C, D; PEPTIDE E-76; CHAIN: X,	THROMBIN LIGHT CHAIN; CHAIN: A, B, C, D; THROMBIN HEAVY CHAIN; CHAIN: M, O, P; THROMBOMODULIN; CHAIN: I, I, K, L; THROMBIN INHIBITOR L- GLU-L-GLY-L-ARM; CHAIN: E, F, G, H;	THROMBIN LIGHT CHAIN; CHAIN: A, B, C, D; THROMBIN HEAVY CHAIN; CHAIN: M, N, O, P; THROMBOMODULIN; CHAIN: I, I, K, L; THROMBIN INHIBITOR L- GLU-L-GLY-L-ARM; CHAIN: E, F, G, H;	THROMBIN LIGHT CHAIN; CHAIN: A, B, C, D; THROMBIN HEAVY CHAIN; CHAIN: M, N, O, P; THROMBOMODULIN; CHAIN: I, J, K, I.; THROMBIN INHIBITOR L- GLU-L-GLY-L-ARM; CHAIN: E, F, G, H;	FIBRILLIN; CHAIN: NULL;
SeqFold Score					
PMF Score		0.51	0.07	0.17	-0.07
Verify Score		-0.14	0.16	0.47	-0.00
PSI BLAST Score		1.2e-17	5.1e-15	5.4e-20	1.2e-17
End AA		221	257	262	216
Start AA		106	151	151	141
Chain ID			-	I	
PDB ID		1dx5	1dx5	1dx5	lemn
SEQ ID NO:		351	351	351	351

PDB annotation	MATRIX, CALCIUM-BINDING, GLYCOPROTEIN, 2 REPEAT, SIGNAL, MULTIGENE FAMILY, DISEASE MUTATION, 3 EGF-LIKE DOMAIN, HUMAN FIBRILLIN-1 FRAGMENT, MATRIX PROTEIN	MATRIX PROTEIN EXTRACELLULAR MATRIX, CALCIUM-BINDING, GLYCOPROTEIN, 2 REPEAT, SIGNAL, MULTIGENE FAMILY, DISEASE MUTATION, 3 EGF-LIKE DOMAIN, HUMAN FIBRILLIN-1 FRAGMENT, MATRIX PROTEIN	SIGNALLING PROTEIN BINDING PROTEIN, CYTOKINE, SIGNALLING PROTEIN	SIGNALLING PROTEIN BINDING PROTEIN, CYTOKINE, SIGNALLING PROTEIN	HYDROLASE PROTEIN-INHIBITOR COMPLEX	BLOOD CLOTTING COMPLEX(SERINE PROTEASE/COFACTOR/LIGAND), BLOOD COAGULATION, 2 SERINE PROTEASE, COMPLEX, CO-FACTOR, RECEPTOR ENZYME, 3 INHIBITOR, GLA, EGF, COMPLEX (SERINE 4 PROTEASE/COFACTOR/LIGAND), BLOOD CLOTTING	GLYCOPROTEIN GLYCOPROTEIN	GLYCOPROTEIN GLYCOPROTEIN STGNATTING PROTEIN TYPE I
Coumpound	·	FIBRILLIN; CHAIN: NULL;	TUMOR NECROSIS FACTOR RECEPTOR; CHAIN: A, B;	TUMOR NECROSIS FACTOR RECEPTOR; CHAIN: A. B;	COAGULATION FACTOR XA; CHAIN: A; COAGULATION FACTOR XA: CHAIN: B:	BLOOD COAGULATION FACTOR VIIA; CHAIN: L; BLOOD COAGULATION FACTOR VIIA; CHAIN: H; SOLUBLE TISSUE FACTOR; CHAIN: T; 5L15; CHAIN: I;	LAMININ; CHAIN: NULL;	LAMININ; CHAIN: NULL; TIMOR NECROSIS
SeqFold Score			77.32					73.83
PMF Score		0.34		0.41	0.16	0.80	-0.12	0.06
Verify Score		0.46		0.53	0.61	0.41	0.11	0.03
PSI BLAST Score		8.5e-20	1.8e-15	1.8e-15	3.6e-13	3.4e-15	1-1	3.4e-13
End		261	286	271	268	273	252	277
Start AA		182	122	123	227	182	109	121 27
Chain ID			А	A	В	ъ		4
PDB CI		lemn	lext	lext	1f0s	l fak	1klo	Incf
SEQ NO.		351	351	351	351	351	351	351

PDB annotation	RECEPTOR, STNFR1; INCF 8 BINDING PROTEIN, CYTOKINE INCF 19	COMPLEX (BLOOD COAGULATION/INHIBITOR) CHRISTMAS FACTOR; COMPLEX, INHIBITOR, HEMOPHILIA/EGF, BLOOD COAGULATION, 2 PLASMA, SERINE PROTEASE, CALCIUM- BINDING, HYDROLASE, 3	COMPLEX (BLOOD COAGULATION/INHIBITOR) CHRISTMAS FACTOR; COMPLEX, INHIBITOR, HEMOPHILIA/EGF, BLOOD COAGULATION, 2 PLASMA, SERINE PROTEASE, CALCIUM- BINDING, HYDROLASE, 3 GLYCOPROTEIN	COMPLEX (BLOOD COAGULATION/INHIBITOR) CHRISTMAS FACTOR; COMPLEX, INHIBITOR, HEMOPHILIA/EGF, BLOOD COAGULATION, 2 PLASMA, SERINE PROTEASE, CALCIUM- BINDING, HYDROLASE, 3 GLYCOPROTEIN	COMPLEX (BLOOD COAGULATION/INHIBITOR) CHRISTMAS FACTOR; COMPLEX, INHIBITOR, HEMOPHILIA/EGF, BLOOD COAGULATION, 2 PLASMA, SERINE PROTEASE, CALCIUM- BINDING, HYDROLASE, 3 GLYCOPROTEIN
Coumpound.	FACTOR RECEPTOR; INCF R 4 CHAIN: A, B; INCF 5	ပ်	FACTOR IXA; CHAIN: C, L,; D-PHE-PRO-ARG; CHAIN: I; IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	FACTOR IXA; CHAIN: C, C L,; D-PHE-PRO-ARG; C CHAIN: I; IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	FACTOR IXA; CHAIN: C, C L,; D-PHE-PRO-ARG; C CHAIN: I, B B B S S S
SeqFold Score				77.38	
PMF Score		0.11	0.04		0.13
Verify Score		-0.17	-0.15		0.09
PSI BLAST Score		5.1e-12	5.4e-10	1.6e-20	1.6e-20
End		180	208	283	276
Start		101	103	143	155
Chain ID		1	ьì	-I	_
PDB ID		1pfx	Ipfx	1pfx	Npfx 1
SEQ ID NO:		351	351	351	351

PDB annotation	COMPLEX (BLOOD COAGULATION/INHIBITOR) CHRISTMAS FACTOR; COMPLEX, INHIBITOR, HEMOPHILIA/EGF, BLOOD COAGULATION, 2 PLASMA, SERINE PROTEASE, CALCIUM- BINDING, HYDROLASE, 3 GLYCOPROTEIN	SERINE PROTEASE FVIIA; FVIIA; BLOOD COAGULATION, SERINE PROTEASE	SERINE PROTEASE FVIIA; FVIIA; BLOOD COAGULATION, SERINE PROTEASE	SERINE PROTEASE FVIIA; FVIIA; BLOOD COAGULATION, SERINE PROTEASE	BLOOD COAGULATION FACTOR STUART FACTOR; BLOOD COAGULATION FACTOR, SERINE PROTEINASE, EPIDERMAL 2 GROWTH FACTOR LIKE DOMAIN
Coumpound	FACTOR IXA; CHAIN: C, L.; D-PHE-PRO-ARG; CHAIN: I;	COAGULATION FACTOR VIIA (LIGHT CHAIN); CHAIN: L; COAGULATION FACTOR VIIA (HEAVY CHAIN); CHAIN: H; TRIPEPTIDYL INHIBITOR; CHAIN: C;	COAGULATION FACTOR VIIA (LIGHT CHAIN); CHAIN: L; COAGULATION FACTOR VIIA (HEAVY CHAIN); CHAIN: H; TRIPEPTIDYL INHIBITOR; CHAIN: C;	COAGULATION FACTOR VIIA (LIGHT CHAIN); CHAIN: L; COAGULATION FACTOR VIIA (HEAVY CHAIN); CHAIN: H; TRIPEPTIDYL INHIBITOR; CHAIN: C;	BLOOD COAGULATION FACTOR XA; CHAIN: L, C;
SeqFold Score					
PMF Score	0.93	0.29	0.83	0.93	0.48
Verify Score	0.34	0.13	-0.05	0.52	0.38
PSI BLAST Score	3.46-17	7.2e-20	5.4e-21	1e-13	5.1e-15
End	276	236	263	273	271
Start AA	182	155	185	186	186
Chain ID	1	1	7	1	7
PDB ID	lpfx	1qfk	lqfk	.1qfk	lxka
SEQ ID NO:	351	351	351	351	351

notation																OSOMAL	E, ANTIBIOTIC,	SPECTINOMYCIN,											-	-		
PDB annotation																RIBOSOME 30S RIBOSOMAL	SUBUNIT, RIBOSOME, ANTIBIOTIC,	STREPTOMYCIN, 2 SPECTINOMYCIN,	PAROMOMYCIN													
Coumpound		LECTIN (AGGLUTININ)	WHEAT GERM	AGGLUTININ (ISOLECTIN 2) 9WGA 3	LECTIN (AGGLUTININ)	WHEAT GERM	AGGLUTININ (ISOLECTIN	2) 9WGA 3	LECTIN (AGGLUTININ)	WHEAT GEKIN	AGGLUTININ (ISOLECTIN 2) 9WGA 3	LECTIN (AGGLUTININ)	WHEAT GERM	AGGLUTININ (ISOLECTIN	2) 9WGA 3	16S RIBOSOMAL RNA;	CHAIN: A; FRAGMENT OF	MESSENGER RNA; CHAIN:	X; 30S RIBOSOMAL	PROTEIN S2; CHAIN: B;	30S RIBOSOMAL PROTEIN	S3; CHAIN: C; 30S	RIBOSOMAL PROTEIN S4;	CHAIN: D; 30S	RIBOSOMAL PROTEIN S5;	CHAIN: E; 30S	RIBOSOMAL PROTEIN S6;	CHAIN: F; 30S	RIBOSOMAL PROTEIN S7;	CHAIN: G; 30S	RIBOSOMAL PROTEIN S8;	CHAIN: H; 30S
SeqFold	31030																													_	_	
PMF	3000	-0.13			0.00				0.40			-0.06				0.18																
Verify		0.21			0.04				0.07		-	0.05				-0.06																
PSI BLAST	Score	5.4e-15			1e-18				5.1e-24			1.2e-17				6.8e-44																
End		250			291				200			160				263														_		
Start		109			124				42			6				101																
Chain 1D		А			Ą				⋖			4				3																
PDB CD	1	9wga			9wga				9wga			9wga				lfjg																
SEQ ID	0 0 0	351			351				351			351				355																

PDB annotation																																
Coumpound		RIBOSOMAL PROTEIN S9; CHAIN: 1; 30S RIBOSOMAL	PROTEIN S10; CHAIN: J;	S11; CHAIN: K; 30S	RIBOSOMAL PROTEIN S12;	CHAIN: L; 30S	RIBOSOMAL PROTEIN S13;	CHAIN: M; 30S	RIBOSOMAL PROTEIN S14;	CHAIN: N; 30S	RIBOSOMAL PROTEIN S15;	CHAIN: 0; 30S	RIBOSOMAL PROTEIN S16;	CHAIN: P; 30S	RIBOSOMAL PROTEIN S17;	CHAIN: Q; 30S	RIBOSOMAL PROTEIN S18;	CHAIN: R; 30S	RIBOSOMAL PROTEIN S19;	CHAIN: S; 30S	RIBOSOMAL PROTEIN S20;	CHAIN: T; 30S	RIBOSOMAL PROTEIN	THX; CHAIN: V	RIBOSOMAL PROTEIN	RIBOSOMAL PROTEIN S5	(PROKARYOTIC) 1PKP 3	RIBOSOMAL PROTEIN	RIBOSOMAL PROTEIN SS	(PROKARYOTIC) 1PKP 3	OXIDOREDUCTASE (/NAD\$(D))	((~) ~
SeqFold	Score																											51.18			477.74	
PMF	Score																							,	0.19							
Verify	Score																	-					-		0.37							
PSI	BLAST Score																								1.4e-49			1.4e-49			0	7
End	AA_													_											253			253			337	$\left] \right]$
Start				_																					86			86			7	
Chain	E																									_					~	
PDB	ID																								Ipkp			1pkp			3gpd	
SEQ	e ë											_];	355			355			357	

	T		<u> </u>	Т	1	r			
PDB annotation					TRANSCRIPTION REGULATION PROTO-ONCOGENE, NUCLEAR BODIES (PODS), LEUKEMIA, 2 TRANSCRIPTION REGULATION			TRANSPORT PROTEIN SERINE-RICH RNA POLYMERASE I SUPPRESSOR PROTEIN; ARM REPEAT	LIGASE CBL, UBCH7, ZAP-70, E2, UBIQUITIN, E3, PHOSPHORYLATION, 2 TYROSINE KINASE, UBIQUITINATION, PROTEIN DEGRADATION,
Coumpound		D-GLYCERALDEHYDE-3- PHOSPHATE DEHYDROGENASE (E.C.1.2.1.12) 3GPD 4	OXIDOREDÚCTASE (NAD\$(A)-ALDEHYDE(D)) D-GLYCERALDEHYDE-3- PHOSPHATE DEHYDROGENASE (E.C.1.2.1.12) 3GPD 4		TRANSCRIPTION FACTOR PML; CHAIN: NULL;	VIRUS EQUINE HERPES VIRUS-1 (C3HC4, OR RING DOMAIN) 1CHC 3 (NMR, 1 STRUCTURE) 1CHC 4	VIRUS EQUINE HERPES VIRUS-1 (C3HC4, OR RING DOMAIN) ICHC 3 (NMR, 1 STRUCTURE) 1CHC 4	KARYOPHERIN ALPHA; CHAIN: A, B; MYC PROTO- ONCOGENE PROTEIN; CHAIN: C, D, E, F;	SIGNAL TRANSDUCTION PROTEIN CBL; CHAIN: A; ZAP-70 PEPTIDE; CHAIN: B; UBIQUITIN-CONJUGATING ENZYME E12-18 KDA UBCH7; CHAIN: C;
SeqFold Score									
PMF Score			1.00		0.21	0.31	0.28	0.28	0.04
Verify Score			0.90		-0.73	-0.18	-0.24	90.0	-0.87
PSI BLAST	Score		0		1.8e-09	1.8e-07	0.00034	1.6e-05	5.4e-07
End AA			337		568	267	995	328	565
Start AA			m		524	524	527	188	492
Chain ID			ж					А	∢
PDB ID			3gpd		Ibor	1chc	1chc	lee4	1fbv
SEQ ID	Ö		357		359	359	359	359	359

SEQ	PDB	Chain		End	ISA	Verify	PMF	SeqFold	Coumpound	PDB annotation
a ö	a	9	AA	AA	BLAST Score	Score	Score	Score		
359	1fbv	4	515	567	1.7e-05	-0.22	0.11		SIGNAL TRANSDUCTION PROTEIN CBL; CHAIN: A; ZAP-70 PEPTIDE; CHAIN: B; UBIQUITIN-CONJUGATING ENZYME E12-18 KDA UBCH7; CHAIN: C:	LIGASE CBL, UBCH7, ZAP-70, E2, UBIQUITIN, E3, PHOSPHORYLATION, 2 TYROSINE KINASE, UBIQUITINATION, PROTEIN DEGRADATION,
359	1g25	V	524	568	3.6e-07	-0.35	0.12		CDK-ACTIVATING KINASE ASSEMBLY FACTOR MATI; CHAIN: A;	METAL BINDING PROTEIN RING FINGER PROTEIN MATI; RING FINGER (C3HC4)
361	lelr	A	945	1046	1e-08	0.24	-0.14		TPRZA-DOMAIN OF HOP; CHAIN: A; HSP90-PEPTIDE MEEVD; CHAIN: B;	CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSP90, 2 PROTEIN BINDING
361	1fch	A	794	1063	1.7e-08	0.07	0.04		PEROXISOMAL TARGETING SIGNAL 1 RECEPTOR; CHAIN: A, B; PTS1-CONTAINING PEPTIDE; CHAIN: C, D;	SIGNALING PROTEIN PEROXISMORE RECEPTOR 1, PTS1-BP, PEROXIN-5, PTS1 PROTEIN-PEPTIDE COMPLEX, TETRATRICOPEPTIDE REPEAT, TPR, 2 HELICAL REPEAT
361	1fch	A	942	1120	1.2e-07	0.32	0.05		PEROXISOMAL TARGETING SIGNAL 1 RECEPTOR; CHAIN: A, B; PTS1-CONTAINING PEPTIDE: CHAIN: C, D.	SIGNALING PROTEIN PEROXISMORE RECEPTOR 1, PTS1-BP, PEROXIN-5, PTS1 PROTEIN-PEPTIDE COMPLEX, TETRATRICOPEPTIDE REPEAT, TPR, 2 HELICAL, REPEAT
361	1fch	4	975	1143	8.5e-11	-0.09	0.05		PEROXISOMAL TARGETING SIGNAL 1 RECEPTOR; CHAIN: A, B; PTSI-CONTAINING PEPTIDE; CHAIN: C, D;	SIGNALING PROTEIN PEROXISMORE RECEPTOR 1, PTS1-BP, PEROXIN-5, PTS1 PROTEIN-PEPTIDE COMPLEX, TETRATRICOPEPTIDE REPEAT, TPR, 2 HELICAL REPEAT
362	2bct		352	908	3.6e-17	-0.11	0.83		BETA-CATENIN; CHAIN: NULL;	STRUCTURAL PROTEIN ARMADILLO REPEAT, BETA-CATENIN, STRUCTURAL PROTEIN

PDB annotation	TRANSMEMBRANE PROTEIN COLICIN, BACTERIOCIN, ION CHANNEL FORMATION, TRANSMEMBRANE 2 PROTEIN	STRUCTURAL PROTEIN TWO REPEATS OF SPECTRIN, ALPHA HELICAL LINKER REGION, 2 2 TANDEM 3-HELIX COILED-COILS, STRUCTURAL PROTEIN	TRANSCRIPTION REGULATION SIGMA70; RNA POLYMERASE SIGMA FACTOR, TRANSCRIPTION REGULATION	COMPLEX (INHIBITOR/NUCLEASE) COMPLEX (INHIBITOR/NUCLEASE), COMPLEX (RI-ANG), HYDROLASE 2 MOLECULAR RECOGNITION, EPITOPE MAPPING, LETICINE-RICH 3 REPRATS	COMPLEX (INHIBITOR/NUCLEASE) COMPLEX (INHIBITOR/NUCLEASE), COMPLEX (RI-ANG), HYDROLASE 2 MOLECULAR RECOGNITION, EPITOPE MAPPING, LEUCINE-RICH 3 REPEATS	COMPLEX (INHIBITOR/NUCLEASE) COMPLEX (INHIBITOR/NUCLEASE), COMPLEX (RI-ANG), HYDROLASE 2 MOLECULAR RECOGNITION, EPITOPE MAPPING, LEUCINE-RICH 3 REPEATS	COMPLEX (INHIBITOR/NUCLEASE) COMPLEX (INHIBITOR/NUCLEASE), COMPLEX (RI-ANG), HYDROLASE 2 MOLECULAR RECOGNITION, EPITOPE MAPPING, LEUCINE-RICH 3 REPEATS
Coumpound	COLICIN IA; CHAIN: NULL;	A. B, C;	RNA POLYMERASE PRIMARY SIGMA FACTOR; CHAIN: NULL;	RIBONUCLEASE INHIBITOR; CHAIN: A, D; ANGIOGENIN; CHAIN: B, E;	RIBONUCLEASE INHIBITOR; CHAIN: A, D; ANGIOGENIN; CHAIN: B, E;	RIBONUCLEASE INHIBITOR; CHAIN: A, D; ANGIOGENIN; CHAIN: B, E;	RIBONUCLEASE INHIBITOR; CHAIN: A, D; ANGIOGENIN; CHAIN: B, E;
SeqFold Score	103.86					70.86	
PMF Score		0.48	0.05	-0.02	0.42		0.39
Verify Score		-0.18	-0.40	90.0	-0.01		0.15
PSI BLAST Score	3.4e-10	1.4e-12	7.2e-09	1.7e-17	1.3e-20	1.8e-46	8.5e-16
End AA	632	251	321	543	395	431	349
Start AA	40	37	71	189	219	25	31
Chain ID		V		¥	A	A	4
PDB ID	lcii	lcun	lsig	la4y	1a4y	la4y	la4y
SEQ.	363	363	363	365	365	365	365

SEQ	BDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ö	Œ	Ω	AA	AA	BLAST Score	Score	Score	Score		
365	la4y	A	74	374	1.8e-46	0.18	0.99		RIBONUCLEASE	COMPLEX (INHIBITOR/NUCLEASE)
							-	•	INHIBITOR; CHAIN: A, D; ANGIOGENIN; CHAIN: B,	COMPLEX (INHIBITOR/NUCLEASE), COMPLEX (RI-ANG), HYDROLASE 2
-								_	வீ	MOLECULAR RECOGNITION, EPITOPE
										MAPPING, LEUCINE-RICH 3 REPEATS
365	la9n	∢	=======================================	246	5.4e-24	0.30	0.77		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEINRNA)
									CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
									CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
3,75				1	1				CHAIN: B, D;	
365	la9n	Ą	128	270	5.4e-27	0.34	0.45		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
				_					CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
				_					CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
	_								CHAIN: B, D;	
365	la9n	٧	175	318	3.6e-25	0.55	0.68		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
									CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
	_								CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
									CHAIN: B, D;	
365	1a9n	V	248	390	1.8e-24	0.31	0.78		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
									CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
									CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
									CHAIN: B, D;	
365	la9n	⋖	596	409	1.3e-18	0.33	0.19		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
									CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
_									CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
]									CHAIN: B, D;	
365	1a9n	∀	8	198	1.8e-16	0.27	0.86		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
									CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
_					-				CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
									CHAIN: B, D;	
365	la9n	ပ	128	273	1.6e-27	0.48	0.59		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
			_						CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
				-					CHAIN: A, C; U2 B"; CHAIN: B. D:	RNA, SNRNP, RIBONUCLEOPROTEIN
365	la9n	၁	175	318	1.8e-25	0.22	0.48		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)

PDB annotation	COMPLEX (NUCLEAR PROTEIN/RNA), RNA, SNRNP,RIBONUCLEOPROTEIN	COMPLEX (NUCLEAR PROTEINRNA) COMPLEX (NUCLEAR PROTEINRNA), RNA, SNRNP,RIBONUCLEOPROTEIN	COMPLEX (NUCLEAR PROTEIN/RNA) COMPLEX (NUCLEAR PROTEIN/RNA), RNA, SNRNP, RIBONUCLEOPROTEIN	CELL ADHESION LEUCINE RICH REPEAT, CALCIUM BINDING, CELL ADHESION	CELL ADHESION LEUCINE RICH REPEAT, CALCIUM BINDING, CELL ADHESION	CELL ADHESION LEUCINE RICH REPEAT, CALCIUM BINDING, CELL ADHESION	CELL ADHESION LEUCINE RICH REPEAT, CALCIUM BINDING, CELL ADHESION	CELL ADHESION LEUCINE RICH REPEAT, CALCIUM BINDING, CELL ADHESION	CELL ADHESION LEUCINE RICH REPEAT, CALCIUM BINDING, CELL ADHESION	TRANSFERASE CRYSTAL STRUCTURE, RAB GERANYLGERANYLTRANSFERASE, 2.0 A 2 RESOLUTION, N- FORMYLMETHIONINE, ALPHA
Coumpound	CHAIN: Q, R; U2 A'; CHAIN: A, C; U2 B"; CHAIN: B, D;	U2 RNA HAIRPIN IV; CHAIN: Q, R; U2 A'; CHAIN: A, C; U2 B"; CHAIN: B, D;	U2 RNA HAIRPIN IV; CHAIN: Q, R; U2 A'; CHAIN: A, C; U2 B"; CHAIN: B, D;	INTERNALIN B; CHAIN: A;	INTERNALIN B; CHAIN: A;	INTERNALIN B; CHAIN: A;	INTERNALIN B; CHAIN: A;	INTERNALIN B; CHAIN: A;	INTERNALIN B; CHAIN: A;	RAB GERANYLGERANYLTRAN SFERASE ALPHA SUBUNIT; CHAIN: A, C; RAB
SeqFold Score										
PMF Score		0.68	0.23	1.00	0.62	1.00	0.76	0.31	0.93	1.00
Verify Score		0.35	0.16	0.50	0.58	0.70	0.48	0.42	0.44	0.59
PSI BLAST Score		1.3e-22	3.6e-17	5.1e-24	8.5e-21	6.8e-24	6.8e-22	8.5e-25	1e-24	1.4e-11
End		412	222	320	152	350	417	224	272	157
Start AA		272	74	141	91	213	216	48	93	48
Chain ID		၁	ပ	Ą	4	V	¥	V	V.	4
PDB ID		1a9n	la9n	90P1	1406	1406	140b	1406	140b	1dce
SEQ NO:		365	365	365	365	365	365	365	365	365

PDB annotation	SUBUNIT, BETA SUBUNIT	TRANSFERASE CRYSTAL STRUCTURE, RAB GERANYLGERANYLTRANSFERASE, 2.0 A 2 RESOLUTION, N. FORMYLMETHIONINE, ALPHA SUBUNIT, BETA SUBUNIT	CONTRACTILE PROTEIN LEUCINE- RICH REPEAT, BETA-BETA-ALPHA CYLINDER, DYNEIN, 2 CHLAMYDOMONAS, FLAGELLA	CONTRACTILE PROTEIN LEUCINE- RICH REPEAT, BETA-BETA-ALPHA CYLINDER, DYNEIN, 2 CHLAMYDOMONAS, FLAGELLA	LIGASE CYCLIN A/CDK2- ASSOCIATED PROTEIN P45; CYCLIN A/CDK2-ASSOCIATED PROTEIN P19; SKP1, SKP2, F-BOX, LRR, LEUCINE- RICH REPEAT, SCF, UBIQUITIN, 2 E3, UBIQUITIN PROTEIN LIGASE	LIGASE CYCLIN A/CDK2- ASSOCIATED PROTEIN P45; CYCLIN A/CDK2-ASSOCIATED PROTEIN P19; SKP1, SKP2, F-BOX, LRR, LEUCINE- RICH REPEAT, SCF, UBIQUITIN, 2 E3, UBIQUITIN PROTEIN LIGASE	LIGASE CYCLIN A/CDK2- ASSOCIATED P45; CYCLIN A/CDK2- ASSOCIATED P19; SKP1, SKP2, F-BOX,
Coumpound	GERANYLGERANYLTRAN SFERASE BETA SUBUNIT; CHAIN: B, D;	RAB GERANYLGERANYLTRAN SFERASE ALPHA SUBUNIT; CHAIN: A, C; RAB GERANYLGERANYLTRAN SFERASE BETA SUBUNIT; CHAIN: B, D;	OUTER ARM DYNEIN; CHAIN: A;	OUTER ARM DYNEIN; CHAIN: A;	SKP2; CHAIN: A, C, E, G, I, K, M, O; SKP1; CHAIN: B, D, F, H, J, L, N, P;	SKP2; CHAIN: A, C, E, G, I, K, M, O; SKP1; CHAIN: B, D, F, H, J, L, N, P;	SKP2; CHAIN: A, C; SKP1; CHAIN: B, D;
SeqFold Score	,						
PMF Score		0.49	0.03	0.15	90.0	-0.18	0.07
Verify Score		0.41	-0.48	0.10	-0.08	0.05	0.21
PSI BLAST Score		6.8e-10	1.7e-13	1.7e-11	8.5e-12	1.5e-15	5.1e-14
End		205	343	223	503	290	246
Start AA		76	187	19	220	48	27
Chain ID		А	٧	Ą	¥	A	¥
PDB ID		ldce	1ds9	Ids9	Ifqv	lfqv	1fs2
SEQ ID NO:		365	365	365	365	365	365

PDB annotation	LRRS, LEUCINE-RICH REPEATS, SCF, 2 UBIQUITIN, E3, UBIQUITIN PROTEIN LIGASE	TRANSCRIPTION RNA IP; RANGAP; GTPASE-ACTIVATING PROTEIN FOR	SPII, GTPASE-ACTIVATING PROTEIN, GAP. RNAIP. RANGAP. I.RR.	LEUCINE- 2 RICH REPEAT PROTEIN,	TWINNING, HEMIHEDRAL TWINNING, 3 MEROHEDRAL	TWINNING, MEROHEDRY	ACETYLATION RNASE INHIBITOR, RIBONUCLEASE/ANGIOGENIN	INHIBITOR ACETYLATION, LEUCINE- RICH REPEATS	ACETYLATION RNASE INHIBITOR,	RIBONUCLEASE/ANGIOGENIN	RICH REPEATS	ACETYLATION RNASE INHIBITOR,	INHIBITOR ACETYLATION, LEUCINE-	KICH KEPEA 1S	ACETYLATION RNASE INHIBITOR, RIBONUCLEASE/ANGIOGENIN	INHIBITOR ACETYLATION, LEUCINE-	RICH REPEATS		TYROSINE KINASE X-LINKED	AGAMMAGLOBULINEMIA, XLA, BTK,	SH3 2 DOMAIN, TRANSFERASE	COMPLEX (ADAPTOR PROTEIN/PEPTIDE) ASH, GROWTH
Coumpound	,	GTPASE-ACTIVATING PROTEIN RNA1 SCHPO;	CHAIN: A, B;				RIBONUCLEASE INHIBITOR; CHAIN: NULL;		RIBONUCLEASE	INHIBITOR; CHAIN: NULL;		RIBONUCLEASE NHIBITOR: CHAIN: NI II I:			RIBONUCLEASE INHIBITOR; CHAIN: NULL;			the common of th	KINASE: CHAIN: NIII.			GRB2; CHAIN: A; SOS; CHAIN: B;
SeqFold Score									81.85													
PMF Score		0.01					-0.05					0.17			1.00			;	0.41			0.92
Verify Score		0.21					0.20					0.03		,;;	0.46			000	0.20			0.26
PSI BLAST Score		1.3e-16					1.7e-22		3.6e-60		-	1.7e-20			3.6e-60			_	3.0e-16			3.6e-18
End AA		206					543		431			395	•	ļ	40 1			966	979			325
Start AA		<i>L</i> 9					189		_			31			8			220	907			272
Chain ID		∢																				∢
PDB ID		lyrg					2bnh		2bnh			2bnh			Zbnh				Iaww			laze
SEQ ID NO:		365					365		365			365		3,7,6	365			250)			367

PDB annotation	FACTOR RECEPTOR-BOUND PROTEIN 2; COMPLEX (ADAPTOR PROTEIN/PEPTIDE), SH3 DOMAIN, 2 GUANINE-NUCLEOTIDE RELEASING FACTOR	COMPLEX (ADAPTOR PROTEIN/PEPTIDE) ASH, GROWTH FACTOR RECEPTOR-BOUND PROTEIN 2; COMPLEX (ADAPTOR PROTEIN/PEPTIDE), SH3 DOMAIN, 2 GUANINE-NUCLEOTIDE RELEASING FACTOR	PHOSPHOTRANSFERASE C-SRC, P60-SRC; SRC, TYROSINE KINASE, PHOSPHORYLATION, SH2, SH3, 2 PHOSPHOTYROSINE, PROTO-ONCOGENE, PHOSPHOTRANSFERASE	COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE) COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE), SH3 DOMAIN	COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE) COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE), SH3 DOMAIN	
Coumpound		GRB2; CHAIN: A; SOS; CHAIN: B;	TYROSINE-PROTEIN KINASE SRC; CHAIN: NULL;	GRB2; CHAIN: A; SOS-1; CHAIN: B;	GRB2; CHAIN: A; SOS-1; CHAIN: B;	SIGNAL TRANSDUCTION PROTEIN GROWTH FACTOR RECEPTOR- BOUND PROTEIN 2 (GRB2, N-TERMINAL 1GBR 3 SH3 DOMAIN) COMPLEXED WITH SOS-A PEPTIDE 1GBR 4 (NMR, 29 STRUCTURES) 1GBR 5
SeqFold Score						
PMF Score		1.00	0.55	0.95	86.0	66:0
Verify Score		0.21	0.16	0.74	-0.00	-0.16
PSI BLAST Score		1.4e-17	1.8e-06	1.8e-19	7.2e-17	1.3e-18
End AA			191	327	53	329
Start AA		2	107	271	2	271
Chain ID		Ą		A	A	V V
PDB ID		laze	Ifmk	1gbq	1gbq	1gbr
SEQ NO:		367	367	367	367	367

														_							-			DAPTOP	NOT TO		DAPTOR		DAPTOR
PDB annotation										_														SIGNAL TO ANSPIRED INCIDENTAL LANGIS	SH2, SH3 1GRI 14		SIGNAL TRANSDUCTION ADAPTOR	SH2, SH3 1GRI 14	SIGNAL TRANSDUCTION ADAPTOR
Coumpound		SIGNAL TRANSDUCTION PROTEIN GROWTH	FACTOR RECEPTOR-	N-TERMINAL 1GBR 3 SH3	DOMAIN) COMPLEXED WITH SOS.A PEPTIDE	1GBR 4 (NMR, 29	STRUCTURES) IGBR 5	ADAPTOR PROTEIN	SH3 GROWTH FACTOR	RECEPTOR-BOUND	PROTEIN 2 (GRB2) 1GFC 3	(C-TERMINAL SH3	DOMAIN) (NMR,	MINIMIZED MEAN	STRUCTURE) 1GFC 4	ADAPTOR PROTEIN	CONTAINING SH2 AND	SH3 GROWTH FACTOR	RECEPTOR-BOUND	PROTEIN 2 (GRB2) 1GFC 3	(C-TERMINAL SH3	DOMAIN) (NMR,	MINIMIZED MEAN	GPOWTH FACTOR BOTHIN	PROTEIN 2: 1GRI 5 CHAIN:	A, B; 1GRI 6	GROWTH FACTOR BOUND	PROTEIN 2; 1GRI 5 CHAIN: A. B: 1GRI 6	GROWTH FACTOR BOUND
SeqFold	Score																							20 00					
PMF	Score	0.94					58,	3.								1.00											0.81		1.00
Verify	Score	0.43					000	0.73								-0.21											0.25		0.01
PSI	Score	3.6e-17						1.86-18					-			1.4e-18								1 30-26	2		1.3e-26		1.7e-16
<u> </u>	AA	191					1,1	3								58								330	3		327		59
5 3	AA	24						<u> </u>								4								101			102		4
Chain	a	∢																						A	:	_	V		A
PDB	a	1gbr					3	Igi I								lgfc								.E	-		1gri		1gri
SEQ	NO.	367					2,52	ે જ								367								367			367		367

_						_		_				_		_								_					
PDB annotation	:	SH2, SH3 1GRJ 14	SIGNAL TRANSDUCTION ADAPTOR SH2, SH3 1GRI 14																								PHOSPHOTRANSFERASE PI3K SH3;
Coumpound		PROTEIN 2; 1GRI 5 CHAIN: A, B; 1GRI 6	GROWTH FACTOR BOUND PROTEIN 2; IGRI 5 CHAIN: A, B; IGRI 6	PHOSPHORIC DIESTER HYDROLASE	PHOSPHOLIPASE C-	(E.C.3.1.4.11) 1HSQ 3 (NMR.	MINIMIZED MEAN STRUCTURE) 1HSO 4	PHOSPHORIC DIESTER	HYDROLASE	PHOSPHOLIPASE C-	GAMMA (SH3 DOMAIN)	(E.C.3.1.4.11) 1HSQ 3 (NMR,	MINIMIZED MEAN	STRUCTURE) THSQ 4	PHOSPHORIC DIESTER	PHOSPHOI IPASE	GAMMA (SH3 DOMAIN)	(E.C.3.1.4.11) 1HSO 3 (NMR	MINIMIZED MEAN	STRUCTURE) 1HSQ 4	PHOSPHORIC DIESTER	HYDROLASE	PHOSPHOLIPASE C.	GAMMA (SH3 DOMAIN)	(E.C.3.1.4.11) 1HSQ 3 (NMR,	MINIMIZED MEAN	PHOSPHATIDYLINOSITOL
SeqFold	Score																								_		
PMF	Score		0.42	1.00				0.55							0.62						1.00						0.25
Verify	Score		0.08	0.17				0.20							-0.30						0.37						0.47
ISd	BLAST Score		1.3e-17	1.3e-16				1.8e-16							0.00017						7.2e-17				-		3.6e-13
Ľ.	AA		155	158				328							333						19						161
<u> </u>	AA		20	101				266	-						270						4						102
Chain	a		А																						_		
PDB	a		Igri	lhsq				lhsq				_			Ihsq						lhsq						1pht
SEQ	a ö		367	367				367							367						367						367

PDB CI	 Chain ID	Start	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
				Score				3-KINASE P85-ALPHA SUBUNIT, IPHT 6 CHAIN:	IPHT 9 PHOSPHATIDYLINOSITOL 3- KINASE, P85-ALPHA SUBUNIT, SH3
1pht		271	342	1.6e-15	-0.02	0.17		PHOSPHATIDYLINOSITOL 3-KINASE P85-ALPHA SUBUNIT; IPHT 6 CHAIN:	PHOSPHOTRANSFERASE PI3K SH3; IPHT 9 PHOSPHATIDYLINOSITOL 3- KINASE, P85-ALPHA SUBUNIT, SH3
1pnj		101	191	1.1e-12	0.42	0.05		PHOSPHOTRANSFERASE PHOSPHATIDYLINOSITOL 3-KINASE (P85-ALPHA SUBUNIT, 1PNJ 3 SH3 DOMAIN) (NMR, MINIMIZED AVERAGE STRUCTURE) 1PNJ 4	DOMAIN IPH I 21
lpwt		101	155	1.6e-18	0.80	0.82		ALPHA SPECTRIN; CHAIN: NULL;	CIRCULAR PERMUTANT PWT; CIRCULAR PERMUTANT, SH3 DOMAIN, CYTOSKELETON
lpwt		4	95	1.6e-18	0.25	1.00		ALPHA SPECTRIN; CHAIN: NULL;	CIRCULAR PERMUTANT PWT; CIRCULAR PERMUTANT, SH3 DOMAIN, CYTOSKELETON
1qkw	Ą	100	155	1.le-18	09.0	09.0		ALPHA II SPECTRIN; CHAIN: A;	CYTOSKELETON CYTOSKELETON, MEMBRANE, SH3 DOMAIN
lqkw	A	569	326	5.4e-18	0.42	0.99		ALPHA II SPECTRIN; CHAIN: A;	CYTOSKELETON CYTOSKELETON, MEMBRANE, SH3 DOMAIN
1qkw	 Ą	4	99	1.8e-18	0.19	0.99		ALPHA II SPECTRIN; CHAIN: A;	CYTOSKELETON CYTOSKELETON, MEMBRANE, SH3 DOMAIN
lqly	 ∢	101	155	1.46-17	0.43	0.87		TYROSINE-PROTEIN KINASE BTK; CHAIN: A;	TYROSINE-PROTEIN KINASE BRUTONS TYROSINE KINASE, B CELL PROGENITOR KINASE, TRANSFERASE, TYROSINE-PROTEIN KINASE, PHOSPHORYLATION, 2 SH3 DOMAIN
1sem	A	101	156	1.4e-17	1.29	1.00		SEM-5; 1SEM 3 CHAIN: A, B; 1SEM 5 10-RESIDUE	SIGNAL TRANSDUCTION PROTEIN SRC-HOMOLOGY 3 (SH3) DOMAIN,

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ë	a	a	AA	AA	BLAST Score	Score	Score	Score		
									PROLINE-RICH PEPTIDE FROM MSOS ISEM 8 CHAIN: C, D ISEM 10	PEPTIDE-BINDING PROTEIN, ISEM 18 2 GUANINE NUCLEOTIDE EXCHANGE FACTOR ISEM 19
367	lsem	٧	271	323	1.8e-17	0.11	1.00		SEM-5; ISEM 3 CHAIN: A,	SIGNAL TRANSDUCTION PROTEIN
									B; ISEM 5 10-RESIDUE	SRC-HOMOLOGY 3 (SH3) DOMAIN,
									PROLINE-RICH PEPTIDE	PEPTIDE-BINDING PROTEIN, ISEM 18
	_								CHAIN: C, D ISEM 10	2 GUANINE NUCLEUTIDE EACHANGE FACTOR ISEM 19
367	Isem	A	4	95	7.2e-18	60.0	1.00		SEM-5; ISEM 3 CHAIN: A,	SIGNAL TRANSDUCTION PROTEIN
									B; 1SEM 5 10-RESIDUE	SRC-HOMOLOGY 3 (SH3) DOMAIN,
									PROLINE-RICH PEPTIDE	PEPTIDE-BINDING PROTEIN, 1SEM 18
									FROM MSOS ISEM 8	2 GUANINE NUCLEOTIDE EXCHANGE
									CHAIN: C, D ISEM 10	FACTOR 1SEM 19
367	1sem	٧	4	99	8.5e-18	0.09	1.00		SEM-5; 1SEM 3 CHAIN: A,	SIGNAL TRANSDUCTION PROTEIN
									B; 1SEM 5 10-RESIDUE	SRC-HOMOLOGY 3 (SH3) DOMAIN,
		_							PROLINE-RICH PEPTIDE	PEPTIDE-BINDING PROTEIN, 1SEM 18
				_					FROM MSOS 1SEM 8	2 GUANINE NUCLEOTIDE EXCHANGE
									CHAIN: C, D ISEM 10	FACTOR 1SEM 19
367	Ituc		114	191	7.2e-16	0.41	0.51		ALPHA-SPECTRIN; CHAIN:	CYTOSKELETON CAPPING PROTEIN,
									NULL;	CALCIUM-BINDING, DUPLICATION,
	_									REPEAT, 2 SH3 DOMAIN,
										CYTOSKELETON
367	1tuc		283	338	1.4e-14	-0.16	0.75		ALPHA-SPECTRIN; CHAIN:	CYTOSKELETON CAPPING PROTEIN,
									NULL;	CALCIUM-BINDING, DUPLICATION,
						 -				REPEAT, 2 SH3 DOMAIN,
267	Aholy		151	156	1.00	5	0.50		THE CHANGE OF THE	CYLOSKELEION
/00	40Ck			CCI	76-17 -	ر. در	65.0		HEMAI OPOIEIIC CELL	TRANSFERASE HCK; SH3, PROTEIN
									NINASE; CHAIN: NOLL;	I YKOSINE KINASE, SIGNAL
				1						IKANSDUCTION, 2 IKANSFEKASE
				1						
368	laww		792	328	3.6e-16	0.20	0.41		BRUTON'S TYROSINE KINASE; CHAIN: NULL;	TRANSFERASE ATK, AMGX1, BPK; TYROSINE KINASE, X-LINKED
										AGAMMAGLOBULÍNEMIA, XLA, BTK,
										SH3 2 DUMAIN, IKANSFEKASE

						T -
PDB annotation	COMPLEX (ADAPTOR PROTEIN/PEPTIDE) ASH, GROWTH FACTOR RECEPTOR-BOUND PROTEIN 2; COMPLEX (ADAPTOR PROTEIN/PEPTIDE), SH3 DOMAIN, 2 GUANINE-NUCLEOTIDE RELEASING FACTOR	COMPLEX (ADAPTOR PROTEIN/PEPTIDE) ASH, GROWTH FACTOR RECEPTOR-BOUND PROTEIN 2; COMPLEX (ADAPTOR PROTEIN/PEPTIDE), SH3 DOMAIN, 2 GUANINE-NUCLEOTIDE RELEASING FACTOR	PHOSPHOTRANSFERASE C-SRC, P60-SRC; SRC, TYROSINE KINASE, PHOSPHORYLATION, SH2, SH3, 2 PHOSPHOTYROSINE, PROTOONCOGENE, PHOSPHOTRANSFERASE	COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE) COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE), SH3 DOMAIN	COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE) COMPLEX (SIGNAL TRANSDUCTION/PEPTIDE), SH3 DOMAIN	
Coumpound	GRB2; CHAIN: A; SOS; CHAIN: B;	GRB2; CHAIN: A; SOS; CHAIN: B;	TYROSINE-PROTEIN KINASE SRC; CHAIN: NULL;	GRB2; CHAIN: A; SOS-1; CHAIN: B;	GRB2; CHAIN: A; SOS-1; CHAIN: B;	SIGNAL TRANSDUCTION PROTEIN GROWTH FACTOR RECEPTOR- BOUND PROTEIN 2 (GRB2, N-TERMINAL IGBR 3 SH3 DOMAIN) COMPLEXED WITH SOS-A PEPTIDE
SeqFold Score						
PMF Score	0.92	1.00	0.55	0.95	86.0	0.99
Verify Score	0.26	0.21	0.16	0.74	-0.00	-0.16
PSI BLAST Score	3.6e-18	1.4e-17	1.8e-06	1.8e-19	7.2e-17	1.3e-18
End AA	325	57	161	327	53	329
Start AA	272	2	107	271	2	271
Chain ID	V	Ą		Ą	٧	A
PDB ID	laze	laze	1 fmk	1gbq	1gbq	1gbr
SEQ ID NO:	368	368	368	368	368	368

PDB annotation					3, 2,	-																				ID SIGNAL TRANSDUCTION ADAPTOR	V: SH2, SH3 1GRI 14	┿	ID SIGNAL TRANSDOCTION ADAPTOR N: SH2, SH3 1GRI 14
Coumpound		1GBR 4 (NMR, 29 STRUCTURES) 1GBR 5	SIGNAL TRANSDUCTION PROTEIN GROWTH	FACTOR RECEPTOR-	BOUND PROTEIN 2 (GRB2, N-TERMINAL 1GBR 3 SH3	DOMAIN) COMPLEXED	WITH SOS-A PEPTIDE	IGBR 4 (NMR, 29 STRIICTIMES) IGBR 5	ADAPTOR PROTEIN	CONTAINING SH2 AND	SH3 GROWTH FACTOR	RECEPTOR-BOUND	PROTEIN 2 (GRB2) 1GFC 3	(C-TERMINAL SH3	DOMAIN) (NMR,	MINIMIZED MEAN	STRUCTURE) 1GFC 4	ADAPTOR PROTEIN	CONTAINING SH2 AND	SH3 GROWTH FACTOR	RECEPTOR-BOUND	PROTEIN 2 (GRB2) 1GFC 3	(C-TERMINAL SH3	DOMAIN) (NMK, MINIMIZED MEAN	STRUCTURE) 1GFC 4	GROWTH FACTOR BOUND	PROTEIN 2; IGRI 5 CHAIN:	A, B; IUKI 0	GROW I H FACTOR BOUND PROTEIN 2; 1GRI 5 CHAIN:
SeqFold	250					-																				99.95			
PMF	300		0.94						5	}								1.00			_							100	0.81
Verify Score			0.43	_					0.73) :								-0.21										30.0	0.23
PSI BLAST	Score		3.6e-17						1 80-18	2								1.4e-18								1.3e-26		1 20 26	1.36-20
End			161						155	:	_					_		28		_						330		207	327
Start			94						101	:								4								101		102	102
Chain ID			¥																							Ą		_	A.
PDB CI			1gbr					<u> </u>	1 ofc	b -								1gfc								1gri]-	ıgı
SEQ ID	NO.		368						368									368								368		340	200

PDB annotation		SIGNAL TRANSDUCTION ADAPTOR SH2, SH3 1GRI 14	SIGNAL TRANSDUCTION ADAPTOR SH2, SH3 IGRI 14																				
Coumpound	A, B; 1GRI 6	GROWTH FACTOR BOUND PROTEIN 2; IGRI 5 CHAIN: A. B: IGRI 6	GROWTH FACTOR BOUND PROTEIN 2; IGRI 5 CHAIN: A, B; IGRI 6	PHOSPHORIC DIESTER HYDROLASE PHOSPHOLIPASE C.	GAMMA (SH3 DOMAIN)	(E.C.3.1.4.11) 1HSQ 3 (NMR, MINIMIZED MEAN	STRUCTURE) 1HSQ 4	PHOSPHORIC DIESTER HYDROLASE	PHOSPHOLIPASE C.	GAMMA (SH3 DOMAIN)	(E.C.3.1.4.11) 1HSQ 3 (NMR,	MINIMIZED MEAN	PHOSPHORIC DIESTER	HYDROLASE	PHOSPHOLIPASE C.	GC3141D1HSO30NR	MINIMIZED MEAN	STRUCTURE) 1HSQ 4	PHOSPHORIC DIESTER	HYDROLASE	PHOSPHOLIPASE C-	GAMMA (SH3 DOMAIN)	(E.C.3.1.4.11) IHSQ 3 (NMR, MINIMIZED MEAN
SeqFold Score																							
PMF Score		1.00	0.42	1.00				0.55					0.62			•			1.00				
Verify Score		10.0	0.08	0.17				0.20					-0.30						0.37				
PSI BLAST Score		1.7e-16	1.3e-17	1.3e-16				1.8e-16				•	0.00017						7.2e-17	•			
End		59	155	158				328					333						61				
Start		4	70	101				266					270						4				
Chain ID		Y	4																				
PDB ID		Igri	1gri	lhsq				Ihsq					1hsq	1					lhsq				
SEQ NO:		368	368	368				368					368						368				

PDB annotation		PHOSPHOTRANSFERASE PI3K SH3;	KINASE, P85-ALPHA SUBUNIT, SH3	DOMAIN 1PHT 21	PHOSPHOTRANSFERASE PI3K SH3;	IPHT 9 PHOSPHATIDYLINOSITOL 3-	KINASE, P85-ALPHA SUBUNIT, SH3 DOMAIN 1PHT 21							CIRCULAR PERMUTANT PWT;	CIRCULAR PERMUTANT, SH3	DOMAIN, CYTOSKELETON	CIRCULAR PERMUTANT PWT;	CIRCULAR PERMUTANT, SH3	DOMAIN, CYTOSKELETON	CYTOSKELETON CYTOSKELETON, MEMBRANE SH3 DOMAIN	CYTOSKELETON CYTOSKELETON.	MEMBRANE, SH3 DOMAIN	CYTOSKELETON CYTOSKELETON,	MEMBRANE, SH3 DOMAIN	TYROSINE-PROTEIN KINASE	BRUTONS TYROSINE KINASE, B CELL	PROGENITOR KINASE,	TRANSFERASE, TYROSINE-PROTEIN	KINASE, PHOSPHORYLATION, 2 SH3 DOMAIN
Coumpound	STRUCTURE) 1HSO 4	PHOSPHATIDYLINOSITOL	:NIX	\neg	ITOL		SUBUNIT; 1PHT 6 CHAIN: 1 NULL; 1PHT 7	┝	PHOSPHATIDYLINOSITOL	3-KINASE (P85-ALPHA	DOMAIN) (NAME	MINIMIZED AVERAGE	STRUCTURE) IPNJ 4	SPECTRIN; CHAIN:	NULL;		SPECTRIN; CHAIN:	NOLL;		ALPHA II SPECTRIN;	SPECTRIN:		PECTRIN;	CHAIN: A;		KINASE BTK; CHAIN: A;	.		<u></u>
SeqFold Score			•																	-									
PMF Score		0.25			0.17			0.05						0.82			1.00			09:0	0.99		0.99		0.87				
Verify Score		0.47			-0.02			0.42						08.0			0.25			09:0	0.42		0.19		0.43				
PSI BLAST Score		3.6e-13			1.6e-15			1.le-12						1.6e-18			1.6e-18			1.le-18	5.4e-18		1.8e-18		1.4e-17				
End AA		191			342			191						155			26			155	326		99		155				
Start AA		102			271			101						<u></u>			4		1	9	269		4		101				
Chain ID																					V		_ V						
PDB ID		1pht] pht			Ipnj						lpwt			lpwt		1	Iqkw	1qkw		lqkw		Iqly				_
SEQ NO:		368			368			368						368			368			308	368		368		368				

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ö	a	1	AA A	ΑA	BLAST Score	Score	Score	Score		
368	lsem	4	101	156	1.4e-17	1.29	1.00		SEM-5; ISEM 3 CHAIN: A, B; ISEM 5 10-RESIDUE PROLINE-RICH PEPTIDE FROM MSOS ISEM 8 CHAIN: C. D. ISEM 10	SIGNAL TRANSDUCTION PROTEIN SRC-HOMOLOGY 3 (SH3) DOMAIN, PEPTIDE-BINDING PROTEIN, ISEM 18 2 GUANINE NUCLEOTIDE EXCHANGE FACTOR ISEM 19
368	1sem	∀	271	323	1.8e-17	0.11	1.00		SEM-5; ISEM 3 CHAIN: A, B; ISEM 5 10-RESIDUE PROLINE-RICH PEPTIDE FROM MSOS ISEM 8 CHAIN: C. D. ISEM 10	SIGNAL TRANSDUCTION PROTEIN SRC-HOMOLOGY 3 (SH3) DOMAIN, PEPTIDE-BINDING PROTEIN, 1SEM 18 2 GUANINE NUCLEOTIDE EXCHANGE FACTOR 1SEM 19
368	lsem	٧	4	56	7.2e-18	60.0	1.00		SEM-5; ISEM 3 CHAIN: A, B; ISEM 5 10-RESIDUE PROLINE-RICH PEPTIDE FROM MSOS ISEM 8 CHAIN: C, D ISEM 10	SIGNAL TRANSDUCTION PROTEIN SRC-HOMOLOGY 3 (SH3) DOMAIN, PEPTIDE-BINDING PROTEIN, ISEM 18 2 GUANINE NUCLEOTIDE EXCHANGE FACTOR ISEM 19
368	lsem	A	4	26	8.5e-18	60.0	1.00		SEM-5; ISEM 3 CHAIN: A, B; ISEM 5 10-RESIDUE PROLINE-RICH PEPTIDE FROM MSOS ISEM 8 CHAIN: C, D ISEM 10	SIGNAL TRANSDUCTION PROTEIN SRC-HOMOLOGY 3 (SH3) DOMAIN, PEPTIDE-BINDING PROTEIN, 1SEM 18 2 GUANINE NUCLEOTIDE EXCHANGE FACTOR 1SEM 19
368	1tuc		114	191	7.2e-16	0.41	0.51		ALPHA-SPECTRIN; CHAIN: NULL;	CYTOSKELETON CAPPING PROTEIN, CALCIUM-BINDING, DUPLICATION, REPEAT, 2 SH3 DOMAIN, CYTOSKELETON
368	1tuc		283	338	1.4e-14	-0.16	0.75		ALPHA-SPECTRIN; CHAIN: NULL;	CYTOSKELETON CAPPING PROTEIN, CALCIUM-BINDING, DUPLICATION, REPEAT, 2 SH3 DOMAIN, CYTOSKELETON
368	4hck		101	155	9e-17	0.19	0.59		HEMATOPOIETIC CELL KINASE; CHAIN: NULL;	TRANSFERASE HCK; SH3, PROTEIN TYROSINE KINASE, SIGNAL TRANSDUCTION, 2 TRANSFERASE
369	lapm	ш	378	681	0			140.90	TRANSFERASE(PHOSPHO TRANSFERASE) \$C-/AMP\$-	

PDB annotation																													
Coumpound		DEPENDENT PROTEIN KINASE (E.C.2.7.1.37)	(\$C/APK\$) IAPM 3 (CATALYTIC SUBUNIT)	ALPHA ISOENZYME	MUTANT WITH SER 139	AI.A (/S139A\$) COMPLEX	WITH THE PEPTIDE IAPM	5 INHIBITOR PKI(5-24)	AND THE DETERGENT	MEGA-8 1APM 6	TRANSFERASE(PHOSPHO	DEPENDENT PROTEIN	KINASE (E.C.2.7.1.37)	(\$C/APK\$) 1APM 3	(CATALYTIC SUBUNIT)	ALPHA ISOENZYME	MUTANT WITH SER 139	1APM 4 REPLACED BY	ALA (/S139A\$) COMPLEX	WITH THE PEPTIDE 1APM	5 INHIBITOR PKI(5-24)	AND THE DETERGENT	PHOSPHOTRANSFERASE	CAMP-DEPENDENT	PROTEIN KINASE	CATALYTIC SUBUNIT	1CMK 3 (E.C.2.7.1.37)	ICMK 4	PHOSPHOTRANSFERASE CAMP-DEPENDENT
SeqFold	Score																						147.30						
PMF	Score										9.1																		1.00
Verify	Score							_			0.46														_				0.29
PSI	Score Score										0												0						0
End	AA —										829												681						678
Start	A A										388												368						388
Chain	3										四												E						ш
PDB	3										lapm												1cmk		-				Icmk
SEQ	βŞ										369								_				369						369

PDB annotation				TRANSFERASE KINASE DOMAIN, AUTOINHIBITORY FRAGMENT, HOMODIMER	TRANSFERASE KINASE DOMAIN, AUTOINHIBITORY FRAGMENT, HOMODIMER	HYDROLASE HYDROLASE, DEPHOSPHORYLATION
Coumpound	PROTEIN KINASE CATALYTIC SUBUNIT ICMK 3 (E.C.2.7.1.37) ICMK 4	TRANSFERASE(PHOSPHO TRANSFERASE) CAMP- DEPENDENT PROTEIN KINASE (E.C.2.7.1.37) (CAPK) ICTP 3 (CATALYTIC SUBUNIT) ICTP 4	TRANSFERASE(PHOSPHO TRANSFERASE) CAMP- DEPENDENT PROTEIN KINASE (E.C.2.7.1.37) (CAPK) 1CTP 3 (CATALYTIC SUBUNIT) 1CTP 4	SERINE/THREONINE- PROTEIN KINASE PAK- ALPHA; CHAIN: A, B; SERINE/THREONINE- PROTEIN KINASE PAK- ALPHA: CHAIN: C. D:	SERINE/THREONINE- PROTEIN KINASE PAK- ALPHA; CHAIN: A, B; SERINE/THREONINE- PROTEIN KINASE PAK- ALPHA; CHAIN: C, D;	PROTEIN TYROSINE PHOSPHATASE 1B; CHAIN: NULL;
SeqFold Score		149.99				
PMF Score			1.00	0.06	1.00	0.28
Verify Score			0.36	-0.16	0.65	0.33
PSI BLAST Score		0	0	3.4e-31	1.4e-95	1e-57
End		681	878	81	678	323
Start AA		374	388	15	386	48
Chain ID		ក	ம	∢	U	
PDB ID		1ctp	lctp	113m	lf3m	la5y
SEQ ID NO:		369	369	369	369	374

	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
	<u>e</u>	<u>a</u>	AA	¥¥	BLAST Score	Score	Score	Score	•	
	1c83	A	48	335	1.2e-62	0.01	0.05		PROTEIN-TYROSINE PHOSPHATASE 1B; CHAIN: A;	HYDROLASE PTP1B; HYDROLASE, PHOSPHORYLATION, LIGAND, INHIBITOR
	ldSr	A	158	348	1.7e-21	0.00	0.45		PHOSPHOINOSITIDE PHOSPHOTASE PTEN; CHAIN: A:	HYDROLASE C2 DOMAIN, PHOSPHOTIDYLINOSITOL, PHOSPHOTASE, HYDROLASE
	1gwz		48	320	6.8e-58			53.88.	SHP-1; CHAIN: NULL;	HYDROLASE PROTEIN-TYROSINE PHOSPHATASE; HYDROLASE, PROTEIN TYROSINE PHOSPHATASE, CATALYTIC DOMAIN, 2 WPD LOOP, SH2 DOMAIN
	Ilar	А	29	367	1.7e-76	0.01	-0.13		LAR; CHAIN: A, B;	HYDROLASE TYROSINE PHOSPHATEASE, LAR PROTEIN
	llar	В	14	316	5.1e-62	-0.00	-0.08		LAR; CHAIN: A, B;	HYDROLASE TYROSINE PHOSPHATEASE, LAR PROTEIN
$\overline{}$	llar	В	9	367	3.4e-73	0.02	-0.05		LAR; CHAIN: A, B;	HYDROLASE TYROSINE PHOSPHATEASE, LAR PROTEIN
	1mkp	:	171	312	5.4e-37			142.92	PYSTI; CHAIN: NULL;	HYDROLASE DUAL SPECIFICITY PHOSPHATASE, MAP KINASE HYDROLASE
	1mkp		174	311	5.4e-37	0.89	1.00		PYSTI; CHAIN: NULL;	HYDROLASE DUAL SPECIFICITY PHOSPHATASE, MAP KINASE HYDROLASE
	lmkp		174	312	8.5e-27	0.81	1.00		PYSTI; CHAIN: NULL;	HYDROLASE DUAL SPECIFICITY PHOSPHATASE, MAP KINASE HYDROLASE
	lrpm	Ą	27	317	1.5e-66	0.19	-0.02		RECEPTOR PROTEIN TYROSINE PHOSPHATASE MU; CHAIN: A, B;	RECEPTOR DI; RECEPTOR, PHOSPHATASE, SIGNAL TRANSDUCTION, ADHESION, 2 HYDROLASE
	lvhr	Ą	150	321	1.5e-20			99.02	HUMAN VHI-RELATED DUAL-SPECIFICITY PHOSPHATASE CHAIN: A, B;	HYDROLASE VHR; HYDROLASE, PROTEIN DUAL-SPECIFICITY PHOSPHATASE

		PTOR,	TEIN	ASE, EIN	ZINC	JINC 4, 2 X	INC 4, 2	INC
PDB annotation	HYDROLASE VHR; HYDROLASE, PROTEIN DUAL-SPECIFICITY PHOSPHATASE	HYDROLASE DI; HYDROLASE, SIGNAL TRANSDUCTION, RECEPTOR, GLYCOPROTEIN, 2 PHOSPHORYLATION. SIGNAL	HYDROLASE YOPSI, YOP2B, PASTEURELLA X, PTP-ASE, PROTEIN TYROSINE PHOSPHATASE, HYDROLASE	TYROSINE PHOSPHATASE SYP, SHPTP-2; TYROSINE PHOSPHATASE, INSULIN SIGNALING, SH2 PROTEIN	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC
Coumpound	HUMAN VHI-RELATED DUAL-SPECIFICITY PHOSPHATASE CHAIN: A, B;	RECEPTOR PROTEIN TYROSINE PHOSPHATASE ALPHA; CHAIN: A, B;	YERSINIA PROTEIN TYROSINE PHOSPHATASE; CHAIN: NULL;	SHP-2; CHAIN: A, B;	QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	DNA; CHAIN: A, B, D, E;
SeqFold Score						:		
PMF Score	1.00	-0.08	0.01	-0.03	0.70	0.34	1.00	1.00
Verify Score	0.81	0.10	-0.35	0.01	0.30	-0.00	0.47	0.15
PSI BLAST	3core 1.5e-20	5.1e-67	1.1e-06	1.7e-61	1.7e-25	1.7e-44	3.4e-51	le-51
End AA	319	318	321	317	286	286	314	342
Start AA	153	23	217	25		184	233	261
Chain ID	A	ď		A	4	ပ	ပ	ပ
PDB ED	lvhr	lyfo	lytn	2shp	laih	Ітеу	Imey	lmey
SEQ	374	374	374	374	375	375	375	375

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ë	a	OI .	AA	ΑA	BLAST Score	Score	Score	Score		
									PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)
375	lmey	ပ	261	343	le-51			100.95	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA
									PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
										CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)
375	lmey	၁	289	348	1.7e-37	0.34	0.94		DNA; CHAIN: A, B, D, E;	COMPLEX (ZINC FINGER/DNA) ZINC
									PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
										CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)
375	1466	٧	185	323	3.4e-31	-0.23	0.01		TFIIIA; CHAIN: A, D; 5S	COMPLEX (TRANSCRIPTION
									RIBOSOMAL RNA GENE;	REGULATION/DNA) COMPLEX
									CHAIN: B, C, E, F;	(TRANSCRIPTION
										REGULATION/DNA), RNA POLYMERASE III 2 TRANSCRIPTION
						_				INITIATION, ZINC FINGER PROTEIN
375	146	∢	203	348	6.8e-32			66.72	TFIIIA; CHAIN: A, D; 5S RIBOSOMAI RNA GENE:	COMPLEX (TRANSCRIPTION REGIT A TIONONA) COMPLEY
									CHAIN: B, C, E, F;	(TRANSCRIPTION
										REGULATION/DNA), RNA
								-		POLYMERASE III, 2 TRANSCRIPTION
375	1tf6	A	213	344	6.8e-32	0.10	0.86		TFIIIA: CHAIN: A D: 58	COMPLEX (TRANSCRIPTION
									RIBOSOMAL RNA GENE;	REGULATION/DNA) COMPLEX
									CHAIN: B, C, E, F;	(TRANSCRIPTION)
										REGULATION/DNA), RNA
										POLYMERASE III, 2 TRANSCRIPTION
					7					INITIATION, ZINC FINGER PROTEIN
3/2	pqnI	ပ	209	314	le-31	0.34	0.94	-	YY1; CHAIN: C; ADENO-	COMPLEX (TRANSCRIPTION
			·		-				ASSOCIATED VIRUS P5	REGULATION/DNA) YING-YANG 1;
									INITIATION ELEMENT	TRANSCRIPTION INITIATION,

PDB annotation	INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN • RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI, GLI, ZINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING
Coumpound	DNA; CHAIN: A, B;	YY I; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS PS INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS PS INITIATOR ELEMENT DNA; CHAIN: A, B;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN
SeqFold Score		85.44			84.81		
PMF Score			1.00	1.00		0.96	0.98
Verify Score			0.18	0.38		0.19	0.32
PSI BLAST Score		9e-55	9e-55	5.1e-34	1.4e-54	1.7e-32	1.4e-54
End AA		343	342	342	344	341	348
Start AA		233	238	241	201	213	236
Chain ID		U	ပ	U	∢	∢	A
PDB ID		lubd	lubd	1ubd	2gli	2gli	2gli
SEQ ID NO:		375	375	375	375	375	375

Chain Si	S T	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound GLII; CHAIN: A; DNA;	PDB annotation PROTEIN/DNA) FIVE-FINGER GLI; GLI,
27 61	19		1 1	0.0013	-0.86	0.01		CHAIN: C, D; UDP-GALACTOSE-4-	ZINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA) ISOMERASE EPIMERASE; UDP-
			(($\neg \neg$				EPIMERASE; CHAIN: NULL;	GALACTOSE, EPIMERASE, ISOMERASE
134	134		ი		0.45	1.00		CD46; CHAIN: A, B, C, D, E, F;	GLYCOPROTEIN MEMBRANE COFACTOR PROTEIN (MCP); VIRUS RECEPTOR, COMPLEMENT COFACTOR, SHORT CONSENSUS REPEAT, 2 SCR, MEASLES VIRUS, GLYCOPROTEIN
A 35 155 5.	155		ر. در	5.4e-30			91.85	CD46; CHAIN: A, B, C, D, E, F;	GLYCOPROTEIN MEMBRANE COFACTOR PROTEIN (MCP); VIRUS RECEPTOR, COMPLEMENT COFACTOR, SHORT CONSENSUS REPEAT, 2 SCR, MEASLES VIRUS, GLYCOPROTEIN
A 35 156 5.1	156		5.1	5.16-29	0.54	1.00		CD46; CHAIN: A, B, C, D, E, F;	GLYCOPROTEIN MEMBRANE COFACTOR PROTEIN (MCP); VIRUS RECEPTOR, COMPLEMENT COFACTOR, SHORT CONSENSUS REPEAT, 2 SCR, MEASLES VIRUS, GLYCOPROTEIN
A 33 154 1.	154		1	1.2e-26	0.12	0.36		COMPLEMENT CONTROL PROTEIN; CHAIN: A;	COMPLEMENT INHIBITOR VCP, SP35; COMPLEMENT, NMR, MODULES, PROTEIN STRUCTURE, VACCINIA VIRUS
A. 96 173 8	173		∞	8.5e-17	0.06	-0.01		COMPLEMENT CONTROL PROTEIN; CHAIN: A;	COMPLEMENT INHIBITOR VCP, SP35; COMPLEMENT, NMR, MODULES, PROTEIN STRUCTURE, VACCINIA VIRUS

PDB annotation			MEMBRANE ADHESION SHORT CONSENSUS REPEAT, SUSHI, COMPLEMENT CONTROL PROTEIN, 2 N-GLYCOSYLATION, MULTI- DOMAIN, MEMBRANE ADHESION	MEMBRANE ADHESION SHORT CONSENSUS REPEAT, SUSHI, COMPLEMENT CONTROL PROTEIN, 2 N-GLYCOSYLATION, MULTI- DOMAIN, MEMBRANE ADHESION	COMPLEMENT INHIBITOR SP35, VCP, VACCINIA VIRUS SP35, COMPLEMENT INHIBITOR, COMPLEMENT MODULE, SCR, SUSHI DOMAIN, 2 MODULE PAIR	COMPLEMENT INHIBITOR SP35, VCP, VACCINIA VIRUS SP35; COMPLEMENT INHIBITOR, COMPLEMENT MODULE, SCR, SUSHI DOMAIN, 2 MODULE PAIR	COMPLEMENT INHIBITOR SP35, VCP, VACCINIA VIRUS SP35;
Coumpound	GLYCOPROTEIN FACTOR H, 15TH AND 16TH C- MODULE PAIR (NMR, MINIMIZED 1HFHA 1 AVERAGED STRUCTURE) 1HFH 4 1HFHA 5	GLYCOPROTEIN FACTOR H, 15TH AND 16TH C. MODULE PAIR (NMR, MINIMIZED 1HFHA 1 AVERAGED STRUCTURE) 1HFH 4 1HFHA 5	HUMAN BETA2- GLYCOPROTEIN I; CHAIN: A;	HUMAN BETA2- GLYCOPROTEIN I; CHAIN: A;	VACCINIA VIRUS COMPLEMENT CONTROL PROTEIN; CHAIN: NULL;	VACCINIA VIRUS COMPLEMENT CONTROL PROTEIN; CHAIN: NULL;	VACCINIA VIRUS COMPLEMENT CONTROL
SeqFold Score		68.71				67.44	
PMF Score	0.43		0.10	0.78	96.0		0.21
Verify Score	0.34		0.01	0.39	0.23		-0.17
PSI BLAST Score	3.4e-24	3.4e-24	3.4e-26	5.1e-31	1.7e-25	1.7e-25	1.7e-14
End AA	153	153	091	161	154	155	172
Start	32	33	7		33	33	96
Chain ID			A	A			
PDB ID	1hfh	1hfh	1qub	1qub	lvvc	Iwc	lvvc
SEQ NO:	380	380	380	380	380	380	380

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ö	e	e	AA	AA	BLAST Score	Score	Score	Score		
									PROTEIN; CHAIN: NULL;	COMPLEMENT INHIBITOR, COMPLEMENT MODULE, SCR, SUSHI DOMAIN, 2 MODULE PAIR
381	ldnl	В	155	404	0.00054	-0.22	0.09		SYNTAXIN BINDING PROTEIN 1; CHAIN: A; SYNTAXIN 14: CHAIN: R:	ENDOCYTOSIS/EXOCYTOSIS NSECI; PROTEIN-PROTEIN COMPLEX, MULTI- STIRINIT
									,	
383	lavi	4	148	341	7.2e-08			64.61	APOLIPOPROTEIN A-I; CHAIN: A, B, C, D;	LIPID TRANSPORT APO A-I; LIPOPROTEIN, LIPID TRANSPORT,
										CHOLESTEROL METABOLISM, 2 ATHEROSCI FROSIS HDI 1 CAT.
										ACTIVATION
383	1bq0		_	9/	1.7e-23			65.24	DNAJ; CHAIN: NULL;	CHAPERONE HSP40; CHAPERONE,
										HEAT SHOCK, PROTEIN FOLDING, DNAK
383	1bq0		c	11	1.7e-23	06.0	1.00		DNAJ; CHAIN: NULL;	CHAPERONE HSP40; CHAPERONE,
										HEAT SHOCK, PROTEIN FOLDING, DNAK
383	lcun	∢	165	354	7.2e-10	0.33	0.30		ALPHA SPECTRIN; CHAIN:	STRUCTURAL PROTEIN TWO
									, y e	HELICAL LINKER REGION, 22
										TANDEM 3-HELIX COILED-COILS,
383	lez3	A	197	306	1.6e-06	0.22	0.00		SYNTAXIN-1A; CHAIN: A,	ENDOCYTOSIS/EXOCYTOSIS
									В, С;	SYNAPTOTAGMIN ASSOCIATED 35
										KDA PROTEIN, P35A, THREE HELIX
383	1hdj		2	26	1.8e-28	0.45	1.00		HUMAN HSP40; CHAIN:	MOLECULAR CHAPERONE HDJ-1:
									NULL;	MOLECULAR CHAPERONE
383	Ihdj		2	92	1.8e-28			68.95	HUMAN HSP40; CHAIN:	MOLECULAR CHAPERONE HDJ-1;
				7	-+				NULL;	MOLECULAR CHAPERONE
383	Ihdj		7		8.5e-23	0.79	9:	_	HUMAN HSP40; CHAIN:	MOLECULAR CHAPERONE HDJ-1;
									NULL;	MOLECULAR CHAPERONE

PDB Chain Start End	Start		End	,	PSI RI ACT	Verify	PMF	SeqFold	Coumpound	PDB annotation
10 AA AA BLASI	AA AA BLASI Score	Score	Score		200	۹.	Score	Score	İ	
19uu A 156 398 3.6e-07	156 398	398		3.6e-07				9:26	HUMAN SKELETAL MUSCLE ALPHA-ACTININ 2; CHAIN: A;	CONTRACTILE PROTEIN TRIPLE- HELIX COILED COIL, CONTRACTILE PROTEIN
1sig 226 386 5.4e-06 -0.07	386 5.4e-06	386 5.4e-06	5.4e-06		-0.07		0.03		RNA POLYMERASE PRIMARY SIGMA	TRANSCRIPTION REGULATION SIGMA70; RNA POLYMERASE SIGMA
									FACTOR; CHAIN: NULL;	FACTOR, TRANSCRIPTION REGULATION
1bq0 3 76 1e-24 0.84	76 16-24	76 16-24	16-24		0.84		00'1		DNAJ; CHAIN: NULL;	CHAPERONE HSP40; CHAPERONE, HEAT SHOCK, PROTEIN FOLDING, DNAK
1cun A 165 354 7.2e-10 0.33	165 354 7.2e-10	354 7.2e-10	7.2e-10		0.33		0:30		ALPHA SPECTRIN; CHAIN: A, B, C;	STRUCTURAL PROTEIN TWO REPEATS OF SPECTRIN, ALPHA
										HELICAL LINKER REGION, 2.2 TANDEM 3-HELIX COILED-COILS,
										STRUCTURAL PROTEIN
lez3 A 197 306 1.6e-06 0.22	197 306 1.6e-06	306 1.6e-06	1.6e-06		0.22		0.00		SYNTAXIN-1A; CHAIN: A,	ENDOCYTOSIS/EXOCYTOSIS
									j,	SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX
					_					BUNDLE
1hdj 2 71 3.4e-23 0.68	71 3.4e-23	71 3.4e-23	3.4e-23		99.0		1.00		HUMAN HSP40; CHAIN: NULL;	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
1hdj 2 76 1.8e-28 0.45	76 1.8e-28	76 1.8e-28	1.8e-28		0.45		1.00		HUMAN HSP40; CHAIN: NULL:	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
lquu A 156 399 1.8e-14	156 399	399		1.8e-14		,		76.90	HUMAN SKELETAL	CONTRACTILE PROTEIN TRIPLE-
							-		MUSCLE ALPHA-ACTININ	HELIX COILED COIL, CONTRACTILE
736 404 7.75 07	70 07 7 700	70 07 7 700	7.00 0.7	_	0.07		100		2; CHAIN: A;	THE ANICORIDATION PROTITION
	404 /.26-0/	404 /.26-0/	/0-97./)))		0.0		KINA POLYMEKASE PRIMARY SIGMA	I KANSCKIPIJON KEGOLAJIJON SIGMAJO: RNA POI VMFRASE SIGMA
									FACTOR; CHAIN: NULL;	FACTOR, TRANSCRIPTION REGULATION
						$\overline{}$				
1avl A 148 341 7.2e-08	148 341	341		7.2e-08		_		64.61	APOLIPOPROTEIN A-I;	LIPID TRANSPORT APO A-I;
			 .						CHAIN: A, B, C, D;	LIPOPROTEIN, LIPID TRANSPORT, CHOLESTEROL METABOLISM, 2

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ë	e	<u>e</u>	AA	ΨΨ	BLAST Score	Score	Score	Score		
										ATHEROSCLEROSIS, HDL, LCAT-ACTIVATION
384	1bq0		1	92	1.7e-23			65.24	DNAJ; CHAIN: NULL;	CHAPERONE HSP40; CHAPERONE, HEAT SHOCK, PROTEIN FOLDING, DNAK
384	1bq0		m	11	1.7e-23	0.90	1.00		DNAJ; CHAIN: NULL;	CHAPERONE HSP40; CHAPERONE, HEAT SHOCK, PROTEIN FOLDING, DNAK
384	1cun	∀	165	354	7.2e-10	0.33	0.30		ALPHA SPECTRIN; CHAIN: A, B, C;	STRUCTURAL PROTEIN TWO REPEATS OF SPECTRIN, ALPHA HELICAL LINKER REGION, 2 2 TANDEM 3-HELIX COILED-COILS, STRUCTURAL PROTEIN
384	lez3	A	197	306	1.6e-06	0.22	0.00		SYNTAXIN-1A; CHAIN: A, B, C;	ENDOCYTOSIS/EXOCYTOSIS SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX BUNDLE
384	1hdj		2	9/	1.8e-28	0.45	1.00		HUMAN HSP40; CHAIN: NULL;	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
384	1hdj		2	9/	1.8e-28			68.95	HUMAN HSP40; CHAIN: NULL;	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
384	1hdj		2	77	8.5e-23	0.79	1.00		HUMAN HSP40; CHAIN: NULL;	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
384	lquu	Ą	156	398	3.6e-07			69.56	HUMAN SKELETAL MUSCLE ALPHA-ACTININ 2; CHAIN: A;	CONTRACTILE PROTEIN TRIPLE- HELIX COILED COIL, CONTRACTILE PROTEIN
384	lsig		226	386	5.4e-06	-0.07	0.03		RNA POLYMERASE PRIMARY SIGMA FACTOR; CHAIN: NULL;	TRANSCRIPTION REGULATION SIGMA70; RNA POLYMERASE SIGMA FACTOR, TRANSCRIPTION REGULATION
384	1bq0		က	9/2	le-24	0.84	1.00		DNAJ; CHAIN: NULL;	CHAPERONE HSP40; CHAPERONE, HEAT SHOCK, PROTEIN FOLDING, DNAK
384	Icun	A	165	354	7.2e-10	0.33	0.30		ALPHA SPECTRIN; CHAIN:	STRUCTURAL PROTEIN TWO

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									A, B, C;	REPEATS OF SPECTRIN, ALPHA HELICAL LINKER REGION, 2.2 TANDEM 3-HELIX COILED-COILS, STRUCTURAL PROTEIN
384	lez3	A	197	306	1.6e-06	0.22	0.00		SYNTAXIN-IA; CHAIN: A, B, C;	ENDOCYTOSIS/EXOCYTOSIS SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX BUNDLE
384	1hdj		2	71	3.4e-23	99.0	1.00		HUMAN HSP40; CHAIN: NULL;	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
384	Ihdj		2	76	1.8e-28	0.45	1.00		HUMAN HSP40; CHAIN: NULL;	MOLECULAR CHAPERONE HDJ-1; MOLECULAR CHAPERONE
384	Iquu	Ą	156	399	1.8e-14			76.90	HUMAN SKELETAL MUSCLE ALPHA-ACTININ 2; CHAIN: A;	CONTRACTILE PROTEIN TRIPLE- HELIX COILED COIL, CONTRACTILE PROTEIN
384	lsig		226	404	7.2e-07	0.07	0.01		RNA POLYMERASE PRIMARY SIGMA FACTOR; CHAIN: NULL;	TRANSCRIPTION REGULATION SIGMA70; RNA POLYMERASE SIGMA FACTOR, TRANSCRIPTION REGULATION
385	1byr	А	131	294	7.2e-22	0.25	0.95		ENDONUCLEASE; CHAIN: A;	ENDONUCLEASE ENDONUCLEASE, PHOSPHODIESTERASE,
385	lbyr	A	131	303	1.7e-21	-0.29	0.47		ENDONUCLEASE; CHAIN: A;	ENDONUCLEASE ENDONUCLEASE, PHOSPHODIESTERASE,
386	lehd	A	35	78	0.0036	69:0	0.19		AGGLUTININ ISOLECTIN VI; CHAIN: A	PLANT PROTEIN TWO HOMOLOGOUS HEVEIN-LIKE DOMAINS
386	lehd	A	43	88	5.4e-05	1.08	60:0		AGGLUTININ ISOLECTIN VI; CHAIN: A	PLANT PROTEIN TWO HOMOLOGOUS HEVEIN-LIKE DOMAINS
386	leis	Y	43	88	3.6e-05	1.00	0.00		AGGLUTININ ISOLECTIN VI/AGGLUTININ ISOLECTIN V; CHAIN: A;	SUGAR BINDING PROTEIN UDA; LECTIN, HEVEIN DOMAIN, UDA, SUPERANTIGEN
386	len2	А	43	88	3.6e-05	1.12	0.00		AGGLUTININ ISOLECTIN I/AGGLUTININ ISOLECTIN	SUGAR BINDING PROTEIN UDA; LECTIN, HEVEIN DOMAIN, UDA,

	EQ	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
B Ö	e	e	AA	AA	BLAST Score	Score	Score	Score		
									V/ CHAIN: A;	SUPERANTIGEN, SACCHARIDE BINDING
7										
388	lciu		236	438	3.6e-08	0.11	-0.19		CYCLODEXTRIN GLYCOSYLTRANSFERASE	GLYCOSIDASE CGTASE; 1CIU 8 THERMOSTABLE 1CIU 14
									ICIU 7	
388	lcwv	Ą	223	437	1.8e-09	0.05	-0.20		INVASIN; CHAIN: A;	STRUCTURAL PROTEIN INTEGRIN- BINDING PROTEIN, INV GENE
388	Icwv	V	265	437	3.6e-11	0.34	-0.18		INVASIN; CHAIN: A;	STRUCTURAL PROTEIN INTEGRIN- BINDING PROTEIN, INV GENE
388	ldnu	B	765	442	5.4e-14	0.14	-0.14		PAPD-LIKE CHAPERONE	CHAPERONE/STRUCTURAL PROTEIN
						_•			FIMC; CHAIN: A, C, E, G, I, K M O: MANNOSE	CHAPERONE ADHESIN DONOR
_						_			SPECIFIC ADHESIN FIMH:	CHAPERONE/STRUCTURAL PROTEIN
\dashv		,							CHAIN: B, D, F, H, J, L, N, P;	
388	pqzI	α	162	232	0.0072	-0.41	0.25		RAB-3A; CHAIN: A;	COMPLEX (GTP-BINDING/EFFECTOR)
									RABPHILIN-3A; CHAIN: B;	RAS-RELATED PROTEIN RAB3A;
	-,									COMPLEX (GIP-BINDING/EFFECTOR),
_								- "		G PROTEIN, EFFECTOR, RABCDR, 2
										SYNAPIIC EXOCYTOSIS, KAB
†	2han	ر	141	80	0.000	17.0			Caronina trial Company	FROIEIN, KABSA, KABPHILIN
		 ر		0 7 7 9	7/00.0	44.0-	15.0		CYC/ DNA DUPLEX; CHAIN: A, B; HEME	COMPLEX (TRANSCRIPTION FACTOR/DNA) UAS CYC7: HAP1.18:
								-	ACTIVATOR PROTEIN;	COMPLEX (TRANSCRIPTION
									CHAIN: C, D;	FACTOR/DNA), ASYMMETRY, 2
								_		TRANSCRIPTIONAL ACTIVATION,
╁	1		2/5		5	\dashv	1			HYPERACTIVE MUTANT
	Zvsg .		C07	4 4 0	5.46-09	0.02	-0.19		VARIANT SURFACE	MEMBRANE PROTEIN VSG VSG,
_									OLI COFNOI EIIN ILI AI	IK I FANOSOME, AN HGENIC
+									1.24; CHAIN: A, B;	VAKIA I ION, MEMBRANE PROTEIN
+-	Idet	ď	403	202	1000	0.70	000	1		10 101 11 m 100 m 11 m 101 01 1
		1	3	3	100.0		77.0		DINA LIGASE; CHAIN: A, B;	LIGASE AMP COMPLEX, NAD+- DEPENDENT

	_																	
PDB annotation		ANTI-ONCOGENE CELL CYCLE, ANTI- ONCOGENE, REPEAT, ANK REPEAT	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETAI; COMPLEX	REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA;	GABPBETA1; COMPLEX (TRANSCRIPTION	REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN. ETS DOMAIN.	ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR	COMPLEX (TRANSCRIPTION REGIT ATTOM/DNA) GARPAL PHA:	GABPBETAI; COMPLEX	REGULATION/DNA), DNA-BINDING, 2	NUCLEAR PROTEIN, ETS DOMAIN, ANKYDIN REPEATS TO ANSCRIPTION	3 FACTOR	COMPLEX (TRANSCRIPTION REGULATION/DNA) GARPALPHA:	GABPBETA1; COMPLEX	(TRANSCRIPTION	REGULATION/DNA), DNA-BINDING, 2	NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION
Coumpound		TUMOR SUPPRESSOR P16INK4A; CHAIN: NULL;	GA BINDING PROTEIN ALPHA; CHAIN: A; GA BINDING PROTEIN BETA 1: CHAIN: B: DNA: CHAIN:	D, E;	GA BINDING PROTEIN ALPHA; CHAIN: A; GA	BINDING PROTEIN BETA 1; CHAIN: B; DNA; CHAIN:	D, E;		GA BINDING PROTEIN ALPHA: CHAIN: A: GA	BINDING PROTEIN BETA	i, ciraliy. B, Divo, ciraliy. D, E;			GA BINDING PROTEIN ALPHA; CHAIN; A; GA	BINDING PROTEIN BETA	1; CHAIN: B; DNA; CHAIN:	D, E;	
SeqFold Score					62.84							<u></u>						
PMF Score		0.87	1.00						1.00					00.1				
Verify Score		-0.02	0.41						-0.16					0.53	-			
PSI BLAST Score		1.4e-19	5.1e-37		5.1e-37				7.2e-32					3.4e-28				
End AA		190	188		188				188					154				
Start AA		47	44		5				∞					6				
Chain ID			ω		В				В					m				
PDB TD		la5e	lawc		lawc				lawc					Iawc				
SEQ NO:		394	394		394				394					394				

WO 02/059260

	PDB Chain ID ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
, ,									3 FACTOR
1 bd8		12	161	5.le-30	0.07	96.0		P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF
15d8		5	161	5.1e-30			53.62	P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF
1bi7	В	47	190	1.2e-20	-0.42	0.00		CYCLIN-DEPENDENT KINASE 6; CHAIN: A; MULTIPLE TUMOR SUPPRESSOR; CHAIN: B;	COMPLEX (KINASE/ANTI- ONCOGENE) CDK6; P16INK4A, MTS1; CYCLIN DEPENDENT KINASE, CYCLIN DEPENDENT KINASE INHIBITORY 2 PROTEIN, CDK, INK4, CELL CYCLE, MULTIPLE TUMOR SUPPRESSOR, 3 MTS1, COMPLEX (KINASE/ANTI-ONCOGENE) HEADER
1blx	Ф	12	191	1.5e-29	0.35	1.00		CYCLIN-DEPENDENT KINASE 6, CHAIN: A; P19INK4D; CHAIN: B;	COMPLEX (INHIBITOR PROTEIN/KINASE) INHIBITOR PROTEIN, CYCLIN-DEPENDENT KINASE, CELL CYCLE 2 CONTROL, ALPHA/BETA, COMPLEX (INHIBITOR PROTEIN/KINASE)
l blx	В	3	161	1.5e-29			55.99	CYCLIN-DEPENDENT KINASE 6; CHAIN: A; P19INK4D; CHAIN: B;	COMPLEX (INHIBITOR PROTEIN/KINASE) INHIBITOR PROTEIN, CYCLIN-DEPENDENT KINASE, CELL CYCLE 2 CONTROL, ALPHA/BETA, COMPLEX (INHIBITOR PROTEIN/KINASE)
1bu9	¥	34	061	1.2e-31			60.78	CYCLIN-DEPENDENT KINASE 6 INHIBITOR; CHAIN: A;	HORMONE/GROWTH FACTOR P18- INK4C; CELL CYCLE INHIBITOR, P18INK4C, TUMOR, SUPPRESSOR, CYCLIN- 2 DEPENDENT KINASE, HORMONE/GROWTH FACTOR
1bu9	4	6	188	1.2e-31	0.09	0.98		CYCLIN-DEPENDENT KINASE 6 INHIBITOR;	HORMONE/GROWTH FACTOR P18- INK4C; CELL CYCLE INHIBITOR,

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
NÖ.	a	a	AA	AA	BLAST Score	Score	Score	Score		
									CHAIN: A;	P18INK4C, TUMOR, SUPPRESSOR, CYCLIN- 2 DEPENDENT KINASE, HORMONE/GROWTH FACTOR
394	149s	∢	33	190	3.6e-25	-0.25	0.59		CYCLIN-DEPENDENT KINASE 4 INHIBITOR B; CHAIN: A:	SIGNALING PROTEIN HELIX-TURN- HELIX, ANKYRIN REPEAT
394	1d9s	A	47	.190	1.2e-20	-0.03	0.72		CYCLIN-DEPENDENT KINASE 4 INHIBITOR B; CHAIN: A;	SIGNALING PROTEIN HELIX-TURN- HELIX, ANKYRIN REPEAT
394	Idcq	Ą	10	169	1.4e-20	0.00	0.76		PYK2-ASSOCIATED PROTEIN BETA; CHAIN: A;	METAL BINDING PROTEIN ZINC- BINDING MODULE, ANK YRIN REPEATS. METAL BINDING PROTEIN
394	Idcq	A	48	161	6.8e-21	0.31	0.98		PYK2-ASSOCIATED PROTEIN BETA; CHAIN: A;	METAL BINDING PROTEIN ZINC- BINDING MODULE, ANKYRIN REPEATS, METAL, BINDING PROTEIN
394	lihb	A	41	190	1.2e-31			62.01	CYCLIN-DEPENDENT KINASE 6 INHIBITOR; CHAIN: A, B;	CELL CYCLE INHIBITOR P18- INK4C(INK6); CELL CYCLE INHIBITOR, P18-INK4C(INK6), ANKYRIN REPEAT, 2 CDK 4/6 INHIBITOR
394	likn	D	2	76	1.7e-17	-0.16	0.78		NF-KAPPA-B P65 SUBUNIT; CHAIN: A; NF- KAPPA-B P50D SUBUNIT; CHAIN: C; I-KAPPA-B- ALPHA; CHAIN: D;	TRANSCRIPTION FACTOR, 165; P50D; TRANSCRIPTION FACTOR, IKB/NFKB COMPLEX
394	lika 1	D	9	161	1.5e-35			54.22	NF-KAPPA-B P65 SUBUNIT; CHAIN: A; NF- KAPPA-B P50D SUBUNIT; CHAIN: C; I-KAPPA-B- ALPHA; CHAIN: D;	TRANSCRIPTION FACTOR, 165; P50D; TRANSCRIPTION FACTOR, IKB/NFKB COMPLEX
394	1myo		10	139	3.4e-19	0.10	9.		MYOTROPHIN; CHAIN: NULL	ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR, ANK-REPEAT
394	Ітуо		41	175	9e-28	-0.44	0.12		MYOTROPHIN; CHAIN: NULL	ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR, ANK-REPEAT

PDB annotation		ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR, ANK-REPEAT	ANK-REPEAT MYOTROPHIN, ACETYLATION NMR ANK-REPEAT	COMPLEX (TRANSCRIPTION	REG/ANK REPEAT) COMPLEX	(TRANSCRIPTION REGULATION/ANK	REPEAT), ANK YRIN 2 REPEAT HELIX	COMPLEX (TRANSCRIPTION	REG/ANK REPEAT) COMPLEX	(TRANSCRIPTION REGULATION/ANK	REPEAT), ANKYRIN 2 REPEAT HELIX	COMPLEX (TRANSCRIPTION	REG/ANK REPEAT) COMPLEX	(TRANSCRIPTION REGULATION/ANK	REPEAT), ANKYRIN 2 REPEAT HELIX	TRANSCRIPTION REGULATION	TRANSCRIPTION REGULATION,	ANKYRIN REPEATS, CELL-CYCLE	TRANSCRIPTION REGULATION	IKANSCKIPTION KEGULATION, ANKYRIN REPEATS CELL-CYCLE	TRANSCRIPTION REGIT ATTON	TRANSCRIPTION REGULATION,	ANKYRIN REPEATS, CELL-CYCLE	COMPLEX (ANTI-	ONCOGENE/ANKYRIN REPEATS)	P53BP2; ANKYRIN REPEATS, SH3, P53,	TUMOR SUPPRESSOR, MULTIGENE 2	FAMILY, NUCLEAR PROTEIN,	PHOSPHORYLATION, DISEASE	MUTATION, 3 POLYMORPHISM,	COMPLEX (ANTI-	ONCOGENE/ANK YRIN REPEATS)
Coumpound		MYOTROPHIN; CHAIN: NULL	MYOTROPHIN; CHAIN; NULL	NF-KAPPA-B P65; CHAIN:	A, C; NF-KAPPA-B P50;	CHAIN: B, D; I-KAPPA-B-	ALPHA; CHAIN: E, F;	NF-KAPPA-B P65; CHAIN:	A, C; NF-KAPPA-B P50;	CHAIN: B, D; I-KAPPA-B-	ALPHA; CHAIN: E, F;	NF-KAPPA-B P65; CHAIN:	A, C; NF-KAPPA-B P50;	CHAIN: B, D; I-KAPPA-B-	ALPHA; CHAIN: E, F;	REGULATORY PROTEIN	SWI6; CHAIN: A, B;		REGULATORY PROTEIN	SWIG; CHAIN: A, B;	REGULATORY PROTEIN	SWI6; CHAIN: A, B;		P53; CHAIN: A; 53BP2;	CHAIN: B;							
SeqFold	Score							68.62						_					•													
PMF	Score	0.49	86.0	0.92								1.00				0.81			0.95		0.58			96'0								
Verify	Score	-0.28	-0.21	-0.35								0.30				-0.26			-0.16		-0.09			-0.19								
PSI	Score	1.5e-22	3.6e-21	1.7e-17				8.5e-36				8.5e-36				3.4e-07			5.1e-19		3.6e-22			6.8e-20								
End	AA	188	151	26				161				187				177			173		175			187								
Start	AA	45	∞	2				4				6				132			8		∞	_		104						_		
Chain	a l			Э				ш				ш				۷_			۷.		Ą			м								
PDB	a	lmyo	lmyo	lnfi		_		Infi				Juli				lsw6			lsw6		1sw6			lycs								
SEQ	ΒÖ	394	394	394				394				394				394			394		394			394								

PDB annotation	COMPLEX (ANTI- ONCOGENE/ANKYRIN REPEATS) P53BP2; ANKYRIN REPEATS, SH3, P53, TUMOR SUPPRESSOR, MULTIGENE 2 FAMILY, NUCLEAR PROTEIN, PHOSPHORYLATION, DISEASE MUTATION, 3 POLYMORPHISM, COMPLEX (ANTI- ONCOGENE/ANKYRIN REPEATS)	LIM DOMAIN CONTAINING PROTEINS LIM DOMAIN CONTAINING PROTEINS, METAL-BINDING PROTEIN, ZINC 2 FINGER	ACTIN-BINDING PROTEIN ACTIN- BINDING PROTEIN, CALCIUM- BINDING, PHOSPHORYLATION	CONTRACTILE LIM DOMAIN, CRP, NMR, MUSCLE DIFFERENTIATION, CONTRACTILE	CONTRACTILE LIM DOMAIN, CRP, NMR, MUSCLE DIFFERENTIATION, CONTRACTILE	STRUCTURAL PROTEIN CALPONIN HOMOLOGY, ACTIN BINDING, STRUCTURAL PROTEIN	STRUCTURAL PROTEIN CALPONIN HOMOLOGY, ACTIN BINDING, STRUCTURAL PROTEIN	ACTIN-BINDING CALPONIN HOMOLOGY (CH) DOMAIN; FILAMENTOUS ACTIN-BINDING DOMAIN, CYTOSKELETON	ACTIN-BINDING CALPONIN
Coumpound	P53; CHAIN: A; 53BP2; CHAIN: B;	QCRP2 (LIMI); CHAIN: NULL;	T-FIMBRIN; CHAIN: NULL;	CRP1; CHAIN: A;	CRP1; CHAIN: A;	UTROPHIN; CHAIN: A, B;	UTROPHIN; CHAIN: A, B;	SPECTRIN BETA CHAIN; CHAIN: A;	SPECTRIN BETA CHAIN;
SeqFold Score									
PMF Score	0.19	0.72	0.93	0.51	0.48	1.00	1.00	1.00	1.00
Verify Score	-0.53	0.26	-0.05	-0.10	0.20	0.79	1.06	0.91	0.97
PSI BLAST Score	7.2e-23	5.4e-13	8.5e-27	1.8e-13	9e-14	le-12	1.8e-26	6.8e-16	1.1e-27
End	190	522	376	519	519	375	379	383	383
Start AA	42	464	138	432	460	275	281	278	281
Chain ID	В			A	¥	A	V	∢	A
PDB ID	lycs	la7i	laoa	1581	1841	15hd	lbhd	16kr	1bkr
SEQ ID NO:	394	403	403	403	403	403	403	403	403

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeaFold	Coumpound	PDB annotation
ID NO:	a	<u>a</u>	AA	AA	BLAST Score	Score	Score	Score		
									CHAIN: A;	HOMOLOGY (CH) DOMAIN; FILAMENTOUS ACTIN-BINDING DOMAIN, CYTOSKELETON
403	lctl		458	519	3.6e-13	0.29	99:0		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS ICTL 15
403	lcxx	A	464	522	7.2e-13	0.45	0.87		CYSTEINE AND GLYCINE- RICH PROTEIN CRP2; CHAIN: A:	SIGNALING PROTEIN LIM DOMAIN CONTAINING PROTEINS, METAL- BINDING PROTEIN
403	ldxx	∢	134	383	1.5e-36	-0.01	0.18		DYSTROPHIN; CHAIN: A, B, C, D;	STRUCTURAL PROTEIN DYSTROPHIN, MUSCULAR DYSTROPHY, CALPONIN HOMOLOGY DOMAIN, 2 ACTIN-BINDING, UTROPHIN
403	liml		466	519	1.6e-12	0.16	0.77		CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NULL;	METAL-BINDING PROTEIN CRIP; METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN
403	lqag	٧	140	382	1.2e-32	0.15	0.83		UTROPHIN ACTIN BINDING REGION; CHAIN: A, B;	STRUCTURAL PROTEIN CALPONIN HOMOLOGY DOMAIN, DOMAIN SWAPPING, ACTIN BINDING, 2 UTROPHIN, DYSTROPHIN, STRUCTURAL PROTEIN
403	1zfo		464	491	3.6e-06	-0.16	0.41		LASP-1; CHAIN: NULL;	METAL-BINDING PROTEIN LIM DOMAIN, ZINC-FINGER, METAL- BINDING PROTEIN
405	leap	В	131	205	0.0036	0.36	0.05		CATALYTIC ANTIBODY 17E8 COMPLEXED WITH PHENYL [1-(1-N- SUCCINYLAMINO)PENTY L] 1EAP 3 PHOSPHONATE 1EAP 4	
408	Ifsu		21	343	3.4e-37	0.27	-0.07		ż	HYDROLASE ARYLSULFATASE B,

PDB annotation	IN ASB, 4-SULFATASE; SULFATASE, GLYCOSAMINOGLYCAN DEGRADATION, HYDROLASE, SIGNAL, 2 GLYCOPROTEIN, LYSOSOME	, C, STRUCTURAL PROTEIN RETINAL S- ANTIGEN, 48 KD PROTEIN, VISUAL	ARRESTIN, DESENSITISATION OF THE VISUAL TRANSDUCTION 2 CASCADE, BINDING TO ACTICATED AND PHOSPHORY A TED RHODOPSIN	 	THE VISUAL TRANSDUCTION 2 CASCADE, BINDING TO ACTICATED	—ŀ	C, STRUCTURAL PROTEIN RETINAL S- ANTIGEN, 48 KD PROTEIN; VISUAL ARRESTIN, DESENSITISATION OF	THE VISUAL TRANSDUCTION 2 CASCADE, BINDING TO ACTICATED	AND FROSTHOR I LATED KHODOPSIN	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER DNA-BINDING PROTEIN		CONTRACTILE LIM DOMAIN, CRP, NMR, MUSCLE DIFFERENTIATION, CONTRACTILE
Coumpound	ACETYLGALACTOSAMIN E-4-SULFATASE; CHAIN: NULL;	ARRESTIN; CHAIN: A, B, C, D;		ARRESTIN; CHAIN: A, B, C, D;		The state of the s	ARRESTIN; CHAIN: A, B, C, D;			QGSR ZINC FINGER PEPTIDE; CHAIN: A; DI IPI EX	OLIGONUCLEOTIDE BINDING SITE; CHAIN: B,	CRP1; CHAIN: A;
SeqFold Score		83.44				, 20 00	78.07					64.99
PMF Score				-0.18						1.00		
Verify Score				-0.00	-					0.34		
PSI BLAST Score		1.8e-47		1.8e-47		1 7. 64	1./e-54			8.5e-28		7.2e-17
End		370		317		2,5	363			18		219
Start AA		02		22		·	×			2		24
Chain ID		Ą		4		-	۵			K		4
PDB ID		lefi		lcfl		140				la!h		1b8t
SEQ NO:		410		410		415	7			412		412

PDB annotation	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX
Coumpound	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; CR	DNA; CHAIN: A, B, D, E; CC CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; CR CR	DNA; CHAIN: A, B, D, E; CO CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; IN CR	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; CR	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; CR	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; CR	DNA; CHAIN: A, B, D, E; CO CONSENSUS ZINC FINGER FIN PROTEIN; CHAIN: C, F, G; IN1 CR
SeqFold Score			99.05				
PMF Score	00.1	1.00		1.00	1.00	00.1	1.00
Verify Score	0.09	-0.27		-0.05	0.11	0.29	0.11
PSI BLAST Score	le-49	1.7e-50	1.7e-50	8.5e-39	1.7e-49	1.7e-46	1e-49
End AA	221	249	250	260	109	81	137
Start AA	140	168	168	196	28	2	99
Chain ID	ပ	U	U	ပ	O	၁	٥
PDB ID	lmey	Imey	lmey	Imey	lmey	lmey	Imey
SEQ ID NO:	412	412	412	412	412	412	412

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PDB annotation	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA	INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER (DNA)	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INTITATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION	KEGULATION/DNA) YING-YANG I;	INTIATOR ELEMENT, YYI, ZINC 2
Соитроина		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		TFIIIA; CHAIN: A, D; 5S	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;				TFIIIA; CHAIN: A, D; 5S	KIBOSOMAL KNA GENE;	CHAIN: B, C, E, F; .				TFIIIA; CHAIN: A, D; 5S	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;				TFIIIA; CHAIN: A, D; 5S	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;				YY1; CHAIN: C; ADENO-	ASSOCIATED VIKUS PS	DNA; CHAIN: A, B;
SeqFold Score																							104.19								
PMF Score		1.00			0.93					;	0.84						1.00												0.94	-	•
Verify Score		0.36			-0.02						-0.17						-0.20											7	0.05		
PSI BLAST Score		1e-49			1.4e-36						5.le-37						3.4e-37						1.8e-77		•		-		1.8e-58		
End AA		165			258					12.	1/4						153						251						249		
Start AA		8			113					ç	67						m					1	84						138		
Chain ID		ပ			A						<						۷					1	Ψ.					,	ပ		
PDB CI		1mey			1466					1+62	0111						1116					29.7	011						pqnI		
S e S		412			412					212	714					ļ	412					;	715						412		

SEQ NO:	PDB ID	Chain ID	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
										FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)
412	Jubd	၁	142	250	1.8e-58			85.17	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1;
									INITIATOR ELEMENT DNA; CHAIN: A, B;	TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2
										FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX
7.7	1	ļ	Ş	3	20,00	000				(TRANSCRIPTION REGULATION/DNA)
714	70n1	ر	148	249	0.8e-35	60:0-	94		YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1:
									INITIATOR ELEMENT	TRANSCRIPTION INITIATION,
									DNA; CHAIN: A, B;	INITIATOR ELEMENT, YY1, ZINC 2
										FINGER PROTEIN, DNA-PROTEIN
										RECOGNITION, 3 COMPLEX
412	Iubd	ပ	36	137	5.1e-35	0.07	1.00		YYI; CHAIN: C; ADENO-	COMPLEX (TRANSCRIPTION
									ASSOCIATED VIRUS P5	REGULATION/DNA) YING-YANG 1;
			_						INITIATOR ELEMENT	TRANSCRIPTION INITIATION,
									DNA; CHAIN: A, B;	INITIATOR ELEMENT, YY1, ZINC 2
										FINGER PROTEIN, DNA-PROTEIN
										RECOGNITION, 3 COMPLEX
412	1hd	C	,	127	12. 41	010	67.0		VACTO CITABLE OF APPRIL	(IRANSCRIPTION REGULATION/DNA)
714	non!	ر ر	n	. /2/	14-27-7	01.0-	70.0		ASSOCIATED VIRIS PS	REGIT ATTON/DNA) YING-YANG 1:
									INITIATOR ELEMENT	TRANSCRIPTION INITIATION.
									DNA; CHAIN: A, B;	INITIATOR ELEMENT, YY1, ZINC 2
			-							FINGER PROTEIN, DNA-PROTEIN
										RECOGNITION, 3 COMPLEX
										(TRANSCRIPTION REGULATION/DNA)
412	Jubd Jubd	ပ	m	 	8.5e-28	-0.32	1.0		YY1; CHAIN: C; ADENO-	COMPLEX (TRANSCRIPTION
									ASSOCIATED VIRUS PS INITIATOR ELEMENT	REGULATION/DNA) YING-YANG I; TRANSCRIPTION INITIATION,

PDB annotation	INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING PROTEINDNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEINDNA)	COMPLEX (DNA-BINDING
Coumpound	DNA; CHAIN: A, B;	YY I; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INTIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN
SeqFold Score							
PMF Score		1.00	0.98	1.00	0.94	0.84	0.52
Verify Score		0.18	0.10	0.31	0.05	-0.13	-0.05
PSI BLAST Score		3.6e-53	1.7e-34	1.7e-35	5.1e-34	3.46-31	1e-29
End		165	109	193	248	108	258
Start		54	∞	92	120	12	148
Chain ID		ပ	ပ	ပ	∀	٧	A
PDB ID		lubd	lubd	lubd	2gli	2gli	2gli
SEQ ID NO:		412	412	412	412	412	412

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
e ö	e	a	ΨV	AA	BLAST Score	Score	Score	Score		
									GLII; CHAIN: A; DNA; CHAIN: C, D;	PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)
412	2gli	Α .	28	167	3.6e-64	0.12	1.00		ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA-
412	2gli	A	56	195	3.6e-75			91.24	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	BINDING PROTEIN/DNA) COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA-
412	2gli	A	56	195	5.4e-71	0.15	1.00		ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	BINDING PROTEIN/DNA) COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA)
412	2gli	4	5	139	9e-43	-0.57	0.80		ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	COMPLEX (DNA-BINDING) PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA)
412	2gli	A	84	250	3.6e-75	-0.15	0.84		ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)
413	1660		319	371	0.00014	-0.89	0.06		DNA-BINDING PROTEIN HUMAN ENHANCER- BINDING PROTEIN MBP-1 MUTANT WITH CYS 11	
	,								1BBO 3 REPLACED BY ABU (C11ABU) (NMR, 60 STRUCTURES) 1BBO 4	
413	3znf		210	239	1.4e-10	0.20	0.76		ZINC FINGER /DNA\$ BINDING DOMAIN ZINC FINGER (/NMR\$) 3ZNF 3	

		3 FACTOR COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION) REGULATION/DNA), DNA-BINDING, 2	GOMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF	GOMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF	SEACTOR COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF ANKYRIN MOTIF ANKYRIN MOTIF ANKYRIN MOTIF	SEACTOR COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF ANKYRIN MOTIF ANKYRIN MOTIF ANKYRIN MOTIF ANKYRIN MOTIF	GOMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF COMPLEX (KINASE/ANTI- ONCOGENE) CDK6; P16INK44, MTS1; CYCLIN DEPENDENT KINASE, CYCLIN DEPENDENT KINASE INHIBITORY 2 PROTEIN, CDK, INK4, CELL CYCLE, MULTIPLE TUMOR SUPPRESSOR, 3 MTS1, COMPLEX KINASE/ANTI-ONCOGENE) HEADER
	3 FACTOR	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPAL GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BI	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALP GABPBETAI; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BIN NUCLEAR PROTEIN, ETS DOM ANKYRIN REPEATS, TRANSCI 3 FACTOR TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALI GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BIN NUCLEAR PROTEIN, ETS DOW ANKYRIN REPEATS, TRANSCI 3 FACTOR TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALI GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BIN NUCLEAR PROTEIN, ETS DOW ANKYRIN REPEATS, TRANSCI 3 FACTOR TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALI GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BIN NUCLEAR PROTEIN, ETS DON ANKYRIN REPEATS, TRANSCI 3 FACTOR TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF TUMOR SUPPRESSOR, CDK4/6 INHIBIT ANKYRIN MOTIF	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPH/ GABPBETAI; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDI NUCLEAR PROTEIN, ETS DOMAI ANKYRIN REPEATS, TRANSCRIP 3 FACTOR TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR ANKYRIN MOTIF TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR ANKYRIN MOTIF COMPLEX (KINASE/ANTI- ONCOGENE) CDK6; P16INK4A, M CYCLIN DEPENDENT KINASE INHIBITORY 2 PROTEIN, CDK, IN CELL CYCLE, MULTIPLE TUMOR SUPPRESSOR, 3 MTS1, COMPLEX (KINASE/ANTI-ONCOGENE) HEA
TOTO							
NG PROTEIN	ALPHA; CHAIN: A; GA BINDING PROTEIN BETA	1; CHAIN: B; DNA; CHAIN: D, E;	I; CHAIN: B; DNA; CHAIN: D, E; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	I; CHAIN: B; DNA; CHAIN: D, E; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	1; CHAIN: B; DNA; CHAIN: D, E; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	1; CHAIN: B; DNA; CHAIN: B, E, E; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL; P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	B; DNA; CHAIN: CDK4/6 C; CHAIN: NULL; CDK4/6 C; CHAIN: NULL; CDK4/6 C; CHAIN: NULL; CDK4/6 C; CHAIN: NULL; CDK4/6 CGK4/6 CGK4/C CGCK4/C CGCCCC CGCCCC CGCCCC CGCCCC CGCCCC CGCCCC CGCCCC CGCCCC CGCC CGCCC CGC
GA BINDING PROTEIN ALPHA: CHAIN: A: GA	BINDING PRO 1; CHAIN: B; D D F:	ร์	P. C., P., P. P. P. P. P. P. P. P. P. P. P. P. P.	P19INK4D CDK4/6 INHIBITOR; CHAN P19INK4D CDK4/6 INHIBITOR; CHAN	P19INK4D CDK4/6 INHIBITOR; CHAIN P19INK4D CDK4/6 INHIBITOR; CHAIN P19INK4D CDK4/6 INHIBITOR; CHAIN	P19INK4D CDK4/6 INHIBITOR; CHAN P19INK4D CDK4/6 INHIBITOR; CHAN P19INK4D CDK4/6 INHIBITOR; CHAN P19INK4D CDK4/6 INHIBITOR; CHAN	P19INK4D CDK4/6 INHIBITOR; CHAIN: NUL. P19INK4D CDK4/6 INHIBITOR; CHAIN: NUL. P19INK4D CDK4/6 INHIBITOR; CHAIN: NUL. P19INK4D CDK4/6 INHIBITOR; CHAIN: NUL. CYCLIN-DEPENDENT KINASE 6; CHAIN: A; MULTIPLE TUMOR SUPPRESSOR; CHAIN: B;
GA J ALP BIN	1; CH		P19I INH	P19I INHI P19I	P191 P191 INH INH INH	1619 1919 1919 1919 1919 1919	P191 P191
0.99			1.00	1.00	1.00	1.00	1.00
0.09			0.31			0	
8.5e-33 (_	1e-31 0		5.4e-35 0			
329 8		268		277 5			
166		135					55 69
В							В
	lawc	1bd8		1bd8	1bd8 1bd8	1bd8 1bd8 1bd8	
NO:	417	417		417	417	417	417 417 417 417

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PDB annotation	PROTEIN/KINASE)	COMPLEX (INHIBITOR PROTEIN/KINASE) INHIBITOR PROTEIN, CYCLIN-DEPENDENT KINASE, CELL CYCLE 2 CONTROL, ALPHA/BETA, COMPLEX (INHIBITOR	COMPLEX (INHIBITOR PROTEIN/KINASE) INHIBITOR PROTEIN, CYCLIN-DEPENDENT KINASE, CELL CYCLE 2 CONTROL, ALPHA/BETA, COMPLEX (INHIBITOR PROTEIN/KINASE)	HORMONE/GROWTH FACTOR P18- INK4C; CELL CYCLE INHIBITOR, P18INK4C, TUMOR, SUPPRESSOR, CYCLIN- 2 DEPENDENT KINASE, HORMONE/GROWTH FACTOR	HORMONE/GROWTH FACTOR P18- INK4C; CELL CYCLE INHIBITOR, P18INK4C, TUMOR, SUPPRESSOR, CYCLIN- 2 DEPENDENT KINASE, HORMONE/GROWTH FACTOR	SIGNALING PROTEIN HELIX-TURN- HELIX, ANKYRIN REPEAT	METAL BINDING PROTEIN ZINC- BINDING MODULE, ANKYRIN REPEATS, METAL BINDING PROTEIN	CELL CYCLE INHIBITOR P18- INK4C(INK6); CELL CYCLE INHIBITOR, P18-INK4C(INK6), ANKYRIN REPEAT, 2 CDK 4/6 INHIBITOR
Coumpound		CYCLIN-DEPENDENT KINASE 6; CHAIN: A; P19INK4D; CHAIN: B;	CYCLIN-DEPENDENT KINASE 6; CHAIN: A; P19INK4D; CHAIN: B;	CYCLIN-DEPENDENT KINASE 6 INHIBITOR; CHAIN: A;	CYCLIN-DEPENDENT KINASE 6 INHIBITOR; CHAIN: A;	CYCLIN-DEPENDENT KINASE 4 INHIBITOR B; CHAIN: A;	PYK2-ASSOCIATED PROTEIN BETA; CHAIN: A;	CYCLIN-DEPENDENT KINASE 6 INHIBITOR; CHAIN: A, B;
SeqFold Score								
PMF Score		1.00	0.78	0.43	0.00	1.00	1.00	1.00
Verify Score		0.48	-0.07	0.39	0.03	0.47	0.48	0.38
PSI BLAST Score		9e-36	1.7e-25	3.4e-33	8.5e-30	1.4e-35	1.1e-31	3.4e-33
End		279	332	268	334	279	262	268
Start AA		150	169	132	166	155	152	132
Chain ID		В	В	٧	4	¥	Ą	٧
PDB JD		16lx	Jblx	1bu9	1bu9	1d9s	1dcq	lihb
SEQ NO:		417	417	417	417	417	417	417

			T								
PDB annotation		CELL CYCLE INHIBITOR P18- INK4C(INK6); CELL CYCLE INHIBITOR, P18-INK4C(INK6), ANKYRIN REPEAT, 2 CDK 4/6 INHIBITOR	TRANSCRIPTION FACTOR P65; P50D; TRANSCRIPTION FACTOR, IKB/NFKB COMPLEX	ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR. ANK-REPEAT	ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR. ANK-REPEAT	ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR. ANK-REPEAT	ANK-REPEAT MYOTROPHIN, ACETYI ATION NAR ANK-REPEAT	ANK-REPEAT MYOTROPHIN, ACETYLATION, NMR. ANK-REPEAT	COMPLEX (TRANSCRIPTION REG/ANK REPEAT) COMPLEX (TRANSCRIPTION REGULATION/ANK REPEAT) ANK VEIN 2 DEPEAT HEI IV	COMPLEX (ANTI- ONCOGENE/ANKYRIN REPEATS) P33BP2; ANKYRIN REPEATS, SH3, P53, TUMOR SUPPRESSOR, MULTIGENE 2 FAMILY, NUCLEAR PROTEIN, PHOSPHORYLATION, DISEASE MUTATION, 3 POLYMORPHISM, COMPLEX (ANTI-	COMPLEX (ANTI-
Coumpound		CYCLIN-DEPENDENT KINASE 6 INHIBITOR; CHAIN: A, B;	NF-KAPPA-B P65 SUBUNIT; CHAIN: A; NF- KAPPA-B P50D SUBUNIT; CHAIN: C; I-KAPPA-B- ALPHA: CHAIN: D:	MYOTROPHIN; CHAIN: NULL	MYOTROPHIN; CHAIN: NULL	MYOTROPHIN; CHAIN: NULL	MYOTROPHIN; CHAIN:	MYOTROPHIN; CHAIN: NULL	NF-KAPPA-B P65; CHAIN: A, C; NF-KAPPA-B P50; CHAIN: B, D; I-KAPPA-B- A1 PHA-CHAIN: F.	P53; CHAIN: A; 53BP2; CHAIN: B;	P53; CHAIN: A; 53BP2;
SeqFold	20016										
PMF	2 1020	0.71	0.11	0.00	0.88	1.00	1.00	0.41	0.21	1.00	-0.19
Verify Score	2020	0.03	-0.13	-0.32	0.10	80.0	0.40	-0.08	-0.08	0.22	0.09
PSI RI AST	Score	1.7e-29	8.5e-36	1.2e-23	1.4e-25	1.4e-27	1.1e-38	6.8e-17	1.4e-35	3.6e-33	8.5e-13
End		333	329	216	249	245	278	314	329	304	48
Start	1	166	127	103	133	551	163	200	126	164	2
Chain		∢	Ω						ធ	m	В
EDB ID		lihb	1 ikn	lmyo	lmyo	lmyo	Imyo	lmyo	lnfi	1ycs	lycs
SEQ	Ö	417	417	417	417	417	417	417	417	417	417

	P53, E 2	9	, Š, ≻,	r EIN		4,	بر ور			L14,	,15P,	, .		땅		MAL	··	123,	
	ONCOGENE/ANKYRIN REPEATS) P53BP2; ANKYRIN REPEATS, SH3, P53, TUMOR SUPPRESSOR, MULTIGENE 2 FAMILY, NUCLEAR PROTEIN, PHOSPHORYLATION, DISEASE MUTATION, 3 POLYMORPHISM, COMPLEX (ANTI- ONCOGENE/ANKYRIN REPEATS)		COAGOLANI, PETILLIC INHIBITORS, CONFORMATIONAL 2 FLEXIBILITY, SERINE PROTEASE INHIBITOR	RIBOSOME 50S RIBOSOMAL PROTEIN L2P, HMAL2, HL4; 50S RIBOSOMAL	; 50S	RIBOSOMAL PROTEIN L4E, HMAL4,	HL6; 50S RIBOSOMAL PROTEIN L5P,	MAL	SC	RIBOSOMAL PROTEIN L14P, HMAL14,	HL27; 50S RIBOSOMAL PROTEIN L15P,	HIMALIS, HLS; SOS KIBOSOMAL PROTEIN I 18P HMAI 18 HI 12: 50S	RIBOSOMAL PROTEIN L18E, HL29,	L19; 50S RIBOSOMAL PROTEIN L19E,	OMAL	PROTEIN L21E, HL31; 50S RIBOSOMAL	PROTEIN L22P, HMAL22, HL23; 50S	RIBOSOMAL PROTEIN L23P, HMAL23,	HL25, L21; 50S KIBOSOMAL PROTEIN L24P, HMAL24, HL16, HL15; 50S
PDB annotation	ONCOGENE/ANKYRIN REPEATS) P53BP2; ANKYRIN REPEATS, SH3 TUMOR SUPPRESSOR, MULTIGEI FAMILY, NUCLEAR PROTEIN, PHOSPHORYLATION, DISEASE MUTATION, 3 POLYMORPHISM, COMPLEX (ANTI-	ANTI-	COAGOLANI, PEPILDIC INHIB CONFORMATIONAL 2 FLEXIBI SERINE PROTEASE INHIBITOR	DSOMA OS RIBC	PROTEIN L3P, HMAL3, HL1; 50S	IN L4E	L PRO	PROTEIN HS6; 50S RIBOSOMAL	PROTEIN L13P, HMAL13; 50S	IN L14	AL PRO	HMAL 13, HLY, 30S KIBOSOMAL PROTEIN I 18P. HMAI 18, HI 12:	IN L18	L PROT	HMAL 19, HL24; 50S RIBOSOMAI	I; 50S R	L22, HI	IN L231	HL25, L21; 50S KIBOSOMAL PR(L24P, HMAL24, HL16, HL15; 50S
РОВ ап	ANKYH YREN F PRESSC CLEAR LATIO 3 POLY NNTI-	ANTI-COAGULANT ANTI-	I, PEPI FIONAL TEASE	OS RIBO HL4; So	, HMA	PROTE	OSOMA	5, 503 R 6; 50S R	P, HM/	PROTE	MOSOS	P. HMA	PROTE	SOMA	24; 50S	E, HL3	P, HIMA	PROTE	S KIBO 74, HL16
	ONCOGENE/ANK P53BP2; ANKYRIN TUMOR SUPPRES FAMILY, NUCLE/ PHOSPHORYLATI MUTATION, 3 POI COMPLEX (ANTI-	COAG	ORMA'	SOME 5	EIN L3	OMAL	SOS RUB	EIN HS	EIN L13	OMAL	50S RI	15, HU	OMAL	OS RIBO	19, HL	EIN [2]	EIN L22	OMAL	L21; 50 HMAL2
	ONCC P53BI TUMC FAMI PHOS MUT/ COMI	ANTI	CONF	RIBOS L2P, F	PROT	RIBO	HL6;	PROT	PROT	RIBO	HL27;	PROT	RIBOS	L19; 5	HMAI	PROT	PROT	RIBOS	H.25, L.24P,
				. 5S	IN L2;	AL	B.	AL 4	Ď.	Z	2	<u>.</u>	Z		Z		z		z
Coumpound		HIRUSTASIN; CHAIN:		23S RRNA; CHAIN: 0; 5S RRNA; CHAIN: 9;	RIBOSOMAL PROTEIN L2;	CHAIN: A; RIBOSOMAL	PROTEIN L3; CHAIN: B;	CHAIN: C; RIBOSOMAL	PROTEIN LS; CHAIN: D;	RIBOSOMAL PROTEIN	L7AE; CHAIN: E;	FROIE F:	RIBOSOMAL PROTEIN	÷	RIBOSOMAL PROTEIN	֔	RIBOSOMAL PROTEIN	I;	KIBOSOMAL PROTEIN L15; CHAIN: J;
Coun	B;	TASIN	,	23S RRNA; CHAII RRNA; CHAIN: 9;	OMAL	V: A; RII	EN 13	S. C. RII	SIN LS;	OMAL	L7AE; CHAIN: E;	KIBOSOMAL FR L10E: CHAIN: F:	OMAL	L13; CHAIN: G;	OMAL	L14; CHAIN: H;	OMAL	LISE; CHAIN: I;	KIBOSOMAL F L15; CHAIN: J;
	CHAIN: B;	HIRUS	NOLL;	23S RF RRNA	RIBOS	CHAIN	PROTE	CHAID	PROTE	RIBOS	L7AE;	LIOE: (RIBOS	L13; C	RIBOS	L14; C	RIBOS	L15E; (KIBOS L15; CI
SeqFold Score																			
PMF Score		0.01		0.00						-		`							
Verify Score		-0.83		-0.33										-					
PSI BLAST Score		0.0054		6.8e-21															
End		237		156		·		-											
Start AA		209		92							-								
Chain ID				-															
PDB ID		1bx7		ĬĬ															
SEQ ID NO:		419		421															

		-	-	_	_	_										_	_	_	_		_	_		_	_					_		
PDB annotation	RIBOSOMAL PROTEIN L24E, HL21/HL22; 50S RIBOSOMAL PROTEIN	L29P, HMAL29, HL33; 50S RIBOSOMAL	PROTEIN L30P, HMAL30, HL20, HL16;	50S RIBOSOMAL PROTEIN L31E, L34,	HL30; 50S RIBOSOMAL PROTEIN L32E,	HLS; 50S KIBOSOMAL PROTEIN L37E,	L39E, HL39E, HL46E: 50S RIBOSOMAL	PROTEIN L44E, LA, HLA; 50S	RIBOSOMAL PROTEIN L6P, HIMAL6,	HL10 RIBOSOME ASSEMBLY, RNA-	RNA, PROTEIN-RNA, PROTEIN-	PROTEIN																				
Coumpound	RIBOSOMAL PROTEIN L18; CHAIN: K;	RIBOSOMAL PROTEIN	L18E; CHAIN: L;	RIBOSOMAL PROTEIN	L19; CHAIN: M;	KIBOSOMAL PROTEIN	RIBOSOMAL PROTEIN	L22; CHAIN: 0;	RIBOSOMAL PROTEIN	L23; CHAIN: P;	RIBOSOMAL PROTEIN	L24; CHAIN: Q;	RIBOSOMAL PROTEIN	L24E; CHAIN: R;	RIBOSOMAL PROTEIN	L29; CHAIN: S;	RIBOSOMAL PROTEIN	L30; CHAIN: T;	RIBOSOMAL PROTEIN	L31E; CHAIN: U;	RIBOSOMAL PROTEIN	L32E; CHAIN: V;	RIBOSOMAL PROTEIN	L37AE; CHAIN: W;	RIBOSOMAL PROTEIN	L37E; CHAIN: X;	RIBOSOMAL PROTEIN	L39E; CHAIN: Y;	RIBOSOMAL PROTEIN	L44E; CHAIN: Z;	RIBOSOMAL PROTEIN L6;	CHAIN: 1;
SeqFold Score																																
PMF Score																															•	
Verify Score																													·			
PSI BLAST Score				_			_																									
End					•														-													
Start AA																																
Chain ID																																
PDB CI																																
SEQ NO:																																

SEQ NO:	PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
431	la7i		104	651	1.3e-16	-0.32	0.93		QCRP2 (LIMI); CHAIN: NULL;	LIM DOMAIN CONTAINING PROTEINS LIM DOMAIN CONTAINING PROTEINS, METAL-BINDING PROTEIN, ZINC 2 FINGER
431	1a 7 i		104	163	5.1e-13	-0.06	0.72		QCRP2 (LIM1); CHAIN: NULL;	LIM DOMAIN CONTAINING PROTEINS LIM DOMAIN CONTAINING PROTEINS, METAL-BINDING PROTEIN, ZINC 2 FINGER
431	1881	A	26	296	3.4e-25			65.67	CRP1; CHAIN: A;	CONTRACTILE LIM DOMAIN, CRP, NMR, MUSCLE DIFFERENTIATION, CONTRACTILE
431	lctl		991	232	8.5e-14	0.01	0.22		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS ICTL 15
431	1ctl		226	294	3.4e-14	-0.14	0.01		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS ICTL 15
431	Ictl		86	159	1.8e-18	-0.19	0.78		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS ICTL 15
431	Icti		66	159	3.4e-14	0.07	0.57		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS ICTL 15
431	lcxx	¥ _	105	159	3.4e-14	-0.14	99'0		CYSTEINE AND GLYCINE- RICH PROTEIN CRP2; CHAIN: A;	SIGNALING PROTEIN LIM DOMAIN CONTAINING PROTEINS, METAL- BINDING PROTEIN
431	lcxx	Ą	106	159	1.3e-16	-0.22	0.59		CYSTEINE AND GLYCINE- RICH PROTEIN CRP2; CHAIN: A;	SIGNALING PROTEIN LIM DOMAIN CONTAINING PROTEINS, METAL- BINDING PROTEIN
431	liml		104	159	1.7e-13	-0.24	0.82		CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NULL;	METAL-BINDING PROTEIN CRIP; METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN
431	limI		106	171	5.4e-17	-0.29	0.07		CYSTEINE RICH	METAL-BINDING PROTEIN CRIP;

		, LIM	CRIP; LIM	CRIP; LIM		R BETA-		R BETA-	R BETA-		PROTEIN, BENASE		STAI, IIT;	AMMA	PROTEIN,			STA1,	AMMA
PDB annotation		METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN	METAL-BINDING PROTEIN CRIP; METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN	METAL-BINDING PROTEIN CRIP METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN		TRANSCRIPTION INHIBITOR BETA- PROPELLER		TRANSCRIPTION INHIBITOR BETA- PROPELLER	TRANSCRIPTION INHIBITOR BETA-PROPELLER		OXIDOREDUCTASE QUINOPROTEIN, SUPERBARREL, DEHYDROGENASE	COMPLEX (GTP-	BINDING/TRANSDUCER) BETA1, TRANSDUCIN BETA SUBUNIT;	GAMMAI, TRANSDUCIN GAMMA	SOBOMII, COMELEA (GIF- BINDING/TRANSDUCER), G PROTEIN,	HETEROTRIMER 2 SIGNAL TRANSPITCTION	COMPLEX (GTP-	BINDING/TRANSDUCER) BETA1,	GAMMAI, TRANSDUCIN GAMMA SUBINIT: COMPLEX (GTP-
		MET	MET	 		PRO		PRO	TRA		SUPI	SO CO	TRA!	GAIN	BINI	HET	S S	BINE	GAM
Coumpound		INTESTINAL PROTEIN; CHAIN: NULL;	CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NITT	CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NULL:		TRANSCRIPTIONAL REPRESSOR TUP1; CHAIN:	A, B, C;	TRANSCRIPTIONAL REPRESSOR TUP1; CHAIN: A B C:	TRANSCRIPTIONAL REPRESSOR TUPI; CHAIN:	A, B, C;	QUINOPROTEIN ETHANOL DEHYDROGENASE; CHAIN: A, B	GT-ALPHA/GI-ALPHA	CHIMERA; CHAIN: A; GT-BETA; CHAIN: B; GT-	GAMMA; CHAIN: G;			GT-ALPHA/GI-ALPHA	CHIMERA; CHAIN: A; GT-	GAMMA; CHAIN: G;
SeqFold	2000											67.04							
PMF	2000		0.18	0.15		0.28		1.00	1.00		0.12						0.98		
Verify	31000		80.0	0.20	-	0.14		69.0	0.70		-0.29						98.0		
PSI RI ACT	Score		1.7e-13	1.8e-17		3.4e-33		· 1.2e-53	le-50		0.0014	5.1e-56					5.1e-46		
End	Ş		230	235		360		296	349		296	378					297		
Start	S		591	165		171		17	41		205	21					23		
Chain						A		A	A		¥	В					В		
BOA III			liml	liml		lerj		1erj	lerj		Ifig	Igot					lgot		
SEQ m	S S		431	431		441		441	441		441	441					441		

Coumpound PDB annotation	BINDING/TRANSDUCER), G PROTEIN, HETEROTRIMER 2 SIGNAL TRANSDUCTION	GT-ALPHA/GI-ALPHA COMPLEX (GTP- CHIMERA; CHAIN: A; GT- BETA; CHAIN: B; GT- GAMMAI; CHAIN: G; SUBUNIT; COMPLEX (GTP- BINDING/TRANSDUCER), G PROTEIN, HETEROTRIMER 2 SIGNAL	ME CD1 OXIDOREDUCTASE ENZYME, DUCTASE; NITRITE REDUCTASE, OXIDOREDUCTASE, DENITRIFICATION, 2 ELECTRON TRANSPORT PER PLASMIC	CYCLOPHILIN A; CHAIN: COMPLEX (ISOMERASE/PEPTIDE) A; PEPTIDE FROM THE COMPLEX (ISOMERASE/PEPTIDE), HIV-1 CAPSID PROTEIN; CYCLOPHILIN A, HIV-1 CAPSID, 2 CHAIN: B:	ä ÷	IAIN: A, B; MEMBRANE PROTEIN CRYSTAL STRUCTURE, MEMBRANE, FERM DOMAIN. TAIL DOMAIN		1
		GT-ALPHA/GI-ALPHA CHIMERA; CHAIN: A; BETA; CHAIN: B; GT- GAMMA; CHAIN: G;	CYTOCHROME CDI NITRITE REDUCTASE; CHAIN: A, B;	CYCLOPHILIN A; CHAI A; PEPTIDE FROM THE HIV-1 CAPSID PROTEIN CHAIN: B:	CYCLOPHILIN A; CHAI A; PEPTIDE FROM THE HIV-1 CAPSID PROTEIN CHAIN: B;	MOESIN; CHAIN: A, B; MOESIN; CHAIN: C, D;	MOESIN; CHAIN: A, B; MOESIN; CHAIN: C, D;	DADIVEL CITABLE
SeqFold Score		,			72.97			
PMF Score		0.80	-0.15	0.92		1.00	1.00	5
Verify Score		0.45	0.12	-0.02		0.43	0.41	0.63
PSI BLAST Score		5.1e-56	1.3e-16	1.7e-47	1.7e-47	8.5e-55	3.6e-88	5 10 SK
End		370	300	113	114	291	290	201
Start		92	42	27	27	15	25	15
Chain ID		æ	₹	¥.	4	¥	∢	4
PDB TD		1 got	1qks	lawq	lawq	lefl	lefi	1007
SEQ ID NO:		141	441	443	443	446	446	446

PDB annotation	ADHESION	CELL ADHESION 3 SUBDOMAINS, CYTOSKELETON, CELL ADHESION	RNA-BINDING PROTEIN/RNA TRA PRE-MRNA; SPLICING REGULATION, RNP DOMAIN, RNA COMPLEX		RNA-BINDING PROTEIN/RNA TRA	PRE-MRNA; SPLICING REGULATION, P RNP DOMAIN, RNA COMPLEX		RNA-BINDING PROTEIN/RNA TRA	PRE-MRNA; SPLICING REGULATION,			IG GENE REGULATION/RNA POLY(A)			* REGULATION/RNA			┪		_		* REGULATION/RNA			
Coumpound		RADIXIN; CHAIN: A;	SXL-LETHAL PROTEIN; CHAIN: A, B; RNA (5'- R(P*GP*UP*UP*GP*UP*UP *! D*! IP*! IP*! IP*! IP*!	CHAIN: P, Q;	SXL-LETHAL PROTEIN;	CHAIN: A, B; RNA (5'- R(P*GP*UP*UP*UP*UP	*UP*UP*UP*UP*UP*U)-	SXL-LETHAL PROTEIN:	CHAIN: A, B; RNA (5'-	R(P*GP*UP*UP*GP*UP*UP	*UP*UP*UP*UP*UP*U)- CHAIN: P. O;	POLYDENYLATE BINDING	PROTEIN 1; CHAIN: A, B,	C, D, E, F, G, H; RNA (5'-	K(*AP*AP*AP*AP*AP*AP*	AP*AP*AP*AP*A)-3');	CHAIN: M, N, O, F, Q, K, O,	1;	POLYDENYLATE BINDING	PROTEIN 1; CHAIN: A, B,	C, D, E, F, G, H; RNA (5'-	R(*AP*AP*AP*AP*AP*	AP*AP*AP*A)-3');	CHAIN: M, N, O, P, Q, R, S,	Ţ,
SeqFold Score								81.68																	
PMF Score		1.00	0.13		1.00							0.24							9						
Verify Score		0.56	0.13		1.11							0.14							0.71						
PSI BLAST Score		7.2e-88	1.7e-29		1.4e-39			1.4e-39				1.7e-29							1./6-31						
End AA		290	311		232			232				316						١	148						
Start AA		25	154		77			11		_		155						,	7						
Chain ID		¥	∢		4			A				A							<						
PDB ID		1gc7	167f		1b7f			1b7f				lcvj		-					[6]	-					
SEQ ID NO:		446	447		447			447				447			_			1,7	Ì						

PDB annotation	GENE REGULATION/RNA POLY(A) BINDING PROTEIN I, PABP I; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM; PROTEIN-RNA COMPLEX, GENE REGULATION/RNA
Coumpound	POLYDENYLATE BINDING PROTEIN I; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*AP*AP* CHAIN: M, N, O, P, Q, R, S, T.	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A);; CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*AP; CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A); CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A); CHAIN: M, N, O, P, Q, R, S,
SeqFold Score	79.42				72.29
PMF Score		1.00	0.74	1.00	
Verify Score		0.78	0.17	0.48	
PSI BLAST	Score 1.2e-38	1.2e-38	3.4e-27	1.5e-28	5.1e-34
End AA	238	236	298	135	218
Start AA	79	08	155	45	79
Chain ID	4	⋖	æ	В	В
PDB ID	lcvj	Icvj	lcvj	lcvj	lcvj
SEQ		447	447	447	447

End PSI Verify AA BLAST Score Score	PMF	1F SeqFold ore Score	Coumpound	PDB annotation
5.1e-34 0.97 1.00	2		T; POLYDENYLATE BINDING PROTEIN I; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP; CHAIN: M, N, O, P, Q, R, S,	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA
1.7e-23 0.25 0.65	ب ن		T; POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP*AP* AP*AP*AP*AP*AP* CHAIN: M, N, O, P, Q, R, S, T:	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA
1.7e-26 0.64 1.00	2		POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A); CHAIN: M, N, O, P, Q, R, S, T;	GENE REGULATIONRNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATIONRNA
6.8e-24 0.25 0.69	<u>&</u>		POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A); CHAIN: M, N, O, P, Q, R, S, T;	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA
1.7e-20 0.18 0.57	7		POLYDENYLATE BINDING PROTEIN I; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP*AP* AP*AP*AP*AP*AP*AP*	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA

SEQ ID	PDB ID	Chain ID	Start AA	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
Ö					Score					
									CHAIN: M, N, O, P, Q, R, S, T;	
447	lcvj	Н	08	207	3.4e-26	0.62	1.00		POLYDENYLATE BINDING PROTEIN I; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A)-3'); CHAIN: M, N, O, P, Q, R, S, T;	GENE REGULATIONRNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATIONRNA
447	1d8z	Ą	76	152	3.4e-20	1.10 .	1.00		HU ANTIGEN C; CHAIN: A;	RNA BINDING PROTEIN RNA- BINDING DOMAIN
447	1fht		79	164	1.4e-18	1.00	0.99		UI SMALL NUCLEAR RIBONUCLEOPROTEIN A; CHAIN: NULL;	RIBONUCLEOPROTEIN UIAI17; RIBONUCLEOPROTEIN, RNP DOMAIN, SPLICEOSOME
447	1fjc	A	7.1	149	3.6e-18	0.71	0.96		NUCLEOLIN RBD2; CHAIN: A;	STRUCTURAL PROTEIN PROTEIN C23; RNP, RBD, RRM, RNA BINDING DOMAIN, NUCLEOLUS
447	lha1		154	311	8.5e-36	0.05	0.43		HNRNP A1; CHAIN: NULL;	NUCLEAR PROTEIN HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1, NUCLEAR PROTEIN, HNRNP, RBD, RRM, RNP, RNA BINDING, 2 RIBONUCLEOPROTEIN
447	Ihai		74	232	1.4e-48	0.97	1.00		HNRNP AI; CHAIN: NULL;	NUCLEAR PROTEIN HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1, NUCLEAR PROTEIN, HNRNP, RBD, RRM, RNP, RNA BINDING, 2 RIBONUCLEOPROTEIN
447	1hd1	¥	154	232	1.7e-19	0.77	0.98		HETEROGENEOUS NUCLEAR. RIBONUCLEOPROTEIN DO; CHAIN: A;	RNA BINDING PROTEIN RNA- BINDING DOMAIN
447	1hd1	¥	8	149	3.4e-20	1.55	1.00		HETEROGENEOUS	RNA BINDING PROTEIN RNA-

	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
	Ð	AA	ΨΨ	BLAST Score	Score	Score	Score		
								NUCLEAR RIBONUCLEOPROTEIN DO; CHAIN: A;	BINDING DOMAIN
	В	80	149	3.6e-19	0.63	1.00		RIBONUCLEOPROTEIN PROTEIN FROM UI SMALL NUCLEAR	
								RIBONUCLEOPROTEIN (SNRNP UI) INRC 3 (N-	
	-							TERMINAL FRAGMENT, RESIDITES 1 - 95) MITTANT	
	_		-					WITH GLN 85 INRC 4 REPLACED BY CYS (Q85C)	
十	4	S	211	5 10 22	910	0.05		INRC 5	THE CHECK STEEL BY STEEL BY STEEL ST
	¢	3		37-21-5	21.	CC:0		BINDING PROTEIN;	C198, HETEROGENEOUS NUCLEAR
								CHAIN: A;	POLYPYRIMIDINE TRACT BINDING
									PROTEIN, RNP, RNA, SPICING, 2 TRANSLATION
2mss	¥	154	232	6.8e-19	0.62	96.0		MUSASHII; CHAIN: A;	RNA BINDING PROTEIN RNA- BINDING DOMAIN
	A	154	317	1.7e-37	0.17	0.41		HETEROGENEOUS	COMPLEX
								NUCLEAR	(RIBONUCLEOPROTEIN/DNA) HNRNP
	•							RIBONUCLEOPROTEIN A1;	A1, UP1; COMPLEX
								CHAIN: A; 12-	(RIBONUCLEOPROTEIN/DNA),
_								NUCLEOI IDE SINGLE- ETP ANDED TEI OMETBIC	HEIEROGENEOUS NUCLEAR 2 PIBONICI EORDOTERI 41
								DNA; CHAIN; B;	MBONOCEEOFNOIEIN AI
<u> </u>	4	73	238	3.4e-49	0.75	1.00		HETEROGENEOUS	COMPLEX
								NUCLEAR	(RIBONUCLEOPROTEIN/DNA) HNRNP
_	•							RIBONUCLEOPROTEIN A1;	A1, UP1; COMPLEX
								CHAIN: A; 12-	(RIBONUCLEOPROTEIN/DNA),
								NUCLEOTIDE SINGLE-	HETEROGENEOUS NUCLEAR 2
								DNA: CHAIN: B:	MBONOCLEOPROI EIN AI

SEQ ID	PDB ID	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF	SeqFold Score	Coumpound	PDB annotation
NO:					Score					
447	3sxI	۷.	154	311	1.4e-26	0.01	0.18		SEX-LETHAL; CHAIN: A, B, C;	RNA BINDING DOMAIN RNA BINDING DOMAIN, RBD, RNA RECOGNITION
										MOTIF, RRM, 2 SPLICING INHIBITOR, TRANSI ATIONAL INHIBITOR SEX 3
									•	DETERMINATION, X CHROMOSOME
777	-		8	;	3	100	,			DOSAGE COMPENSATION
44	3SXI	∢	%	215	5.le-39	0.87	1.00		SEX-LETHAL; CHAIN: A,	RNA BINDING DOMAIN RNA BINDING
									ဘ ်	DOMAIN, RBD, RNA RECOGNITION
							•••			MOLIF, KKM, 2 SPLICING INHIBITOR,
					-					DETERMINATION, X CHROMOSOME
										DOSAGE COMPENSATION
447	3sxl	4	78	223	5.1e-39			78.28	SEX-LETHAL; CHAIN: A,	RNA BINDING DOMAIN RNA BINDING
				-					j, j	DOMAIN, RBD, RNA RECOGNITION
										MOTIF, RRM, 2 SPLICING INHIBITOR,
										TRANSLATIONAL INHIBITOR, SEX 3
										DETERMINATION, X CHROMOSOME DOSAGE COMPENSATION
448	1a09	∢	68	192	3.4e-24			78.11	C-SRC TYROSINE KINASE;	COMPLEX (TRANSFERASE/PEPTIDE)
-									CHAIN: A, B; ACE-	COMPLEX (TRANSFERASE/PEPTIDE)
									GIT-ON-DIBENTY	
									AMINE); CHAIN: C, D;	
448	1a09	٧	93	192	3.4e-24	1.16	1.00		C-SRC TYROSINE KINASE;	COMPLEX (TRANSFERASE/PEPTIDE)
	_			-					CHAIN: A, B; ACE.	COMPLEX (TRANSFERASE/PEPTIDE)
			•			•			GILL-(N.N-DIPENTY)	
									AMINE); CHAIN; C. D.	
448	1a81	Ξ	1	194	3.4e-16			52.85	SYK KINASE; CHAIN: A, C,	COMPLEX (TRANSFERASE/PEPTIDE)
								•	E, G, I, K; T-CELL	ITAM PEPTIDE; COMPLEX
									SURFACE GLYCOPROTEIN	(TRANSFERASE/PEPTIDE), SYK,
			_						CD3 EPSILON CHAIN;	KINASE, SH2 DOMAIN, ITAM
									CHAIN: B, D, F, H, J, L;	

PDB CI	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
1ab2		84	196	Score 1.7e-18			56.12	TRANSFERASE(PHOSPHO TRANSFERASE) PROTO- ONCOGENE TYPOSINE	
								KINASE (E.C.2.7.1.112) 1AB2 3 (SRC HOMOLOGY 2	
								DOMAIN) (ABELSON, SH2 ABL) 1AB2 4 (NMR, 20 STRUCTURES) 1AB2 5	
laot	ഥ	88	193	1.4e-22			77.83	FYN PROTEIN-TYROSINE KINASE: CHAIN: F:	COMPLEX (PROTO- ONCOGENE/EARLY PROTEIN) SRC
								PHOSPHOTYROSYL PEPTINE: CHAIN: D	HOMOLOGY 2 DOMAIN; SH2
									PEPTIDE COMPLEX, 2 COMPLEX (PROTO-ONCOGENE/EARLY
1441		3	100	1 20 26			72 30	The Construction of the Co	PROTEIN)
,		76	3	1.45-43			92.70	KINASE TRANSFORMING	V-SKC SH2 DOMAIN SKC SH2; V-SKC SH2 DOMAIN, PHOSPHOTYROSINE
								PROTEIN; CHAIN: NULL;	RECOGNITION DOMAIN, PP60 2 SRC SH2 DOMAIN
1bkl		93	196	1.2e-25	1.08	1.00		PP60 V-SRC TYROSINE	V-SRC SH2 DOMAIN SRC SH2; V-SRC
							ı	KINASE TRANSFORMING	SH2 DOMAIN, PHOSPHOTYROSINE
								PROTEIN; CHAIN: NULL;	RECOGNITION DOMAIN, PP60 2 SRC SH2 DOMAIN
1blj		81	195	1.4e-23			90.55	P55 BLK PROTEIN	PHOSPHORYLATION SIGNAL
								TYROSINE KINASE;	TRANSDUCTION, TYROSINE KINASE,
								CHAIN: NULL;	TRANSFERASE, 2
									PHOSPHOTRANSFERASE, PHOSPHORYLATION
lcwd	T	93	189	9e-24			91.11	P56LCK TYROSINE	COMPLEX
_					-			KINASE; CHAIN: L;	(PHOSPHOTRANSFERASE/PEPTIDE)
								PHOSPHONOPEPTIDE	PHOSPHOTRANSFERASE, COMPLEX
†								CHAIN: P;	(PHOSPHOTRANSFERASE/PEPTIDE)
lefin	∢	37	&	1.7e-10	-0.0 4	0.03	,	FYN TYROSINE KINASE; CHAIN: A. C. HIV-1 NFF	COMPLEX (SH3 DOMAIN/VIRAL FNHANCER) SRC-HOMOLOGY 3
								CHAIN: A, C; HIV-1 NEF	ENHANCEK) SKC-F

PDB annotation	DOMAN; COMPLEX (SH3 DOMAIN/YIRAL ENHANCER), PROTO- ONCOGENE, 2 TRANSFERASE, TYROSINE-PROTEIN KINASE, PHOSPHORYLATION, 3 AIDS, MYRISTYLATION, GTP-BINDING, ATP-BINDING, SH3 DOMAIN, 4 SH2 DOMAIN, PPII HELIX, PXXP MOTIF	PHOSPHOTRANSFERASE C-SRC, P60-SRC; SRC, TYROSINE KINASE, PHOSPHORYLATION, SH2, SH3, 2 PHOSPHOTYROSINE, PROTO-ONCOGENE, PHOSPHOTYRASINE, PROTO-			SIGNAL TRANSDUCTION ADAPTOR SH2, SH3 1GR1 14
Coumpound	PROTEIN; CHAIN: B, D;	TYROSINE-PROTEIN KINASE SRC; CHAIN: NULL;	SIGNAL TRANSDUCTION PROTEIN GROWTH FACTOR RECEPTOR- BOUND PROTEIN 2 (GRB2, N-TERMINAL 1GBR 3 SH3 DOMAIN) COMPLEXED WITH SOS-A PEPTIDE 1GBR 4 (NMR, 29 STRUCTURES) 1GBR 5	ADAPTOR PROTEIN CONTAINING SH2 AND SH3 GROWTH FACTOR RECEPTOR-BOUND PROTEIN 2 (GRB2) 1GFC 3 (C-TERMINAL SH3 DOMAIN) (NMR, MINIMIZED MEAN STRUCTURE) 1GFC 4	GROWTH FACTOR BOUND PROTEIN 2; 1GRI 5 CHAIN: A, B; 1GRI 6
SeqFold Score		i			50.22
PMF Score		1.00	0.23	-0.11	
Verify Score		09.0	0.01	0.25	
PSI BLAST Score		3.4e-42	3.4e-10	3.46-10	1.5e-23
End AA		207	93	88	209
Start AA		34	27	37	35
Chain ID			∢		A
PDB ID		1fmk	1gbr	1gfc	lgri
SEQ ID NO:		448	448	448	448

pound PDB annotation	TOR BOUND SIGNAL TRANSDUCTION ADAPTOR SRI 5 CHAIN: SH2, SH3 1GRI 14	DIESTER ASE C- DOMAIN) HSQ 3 (NMR, EAN 1HSO 4	TROSINE COMPLEX (KINASE/PEPTIDE) (7 CHAIN: A; TIDE 40)YQPQPA; N: B; ILCK	YROSINE COMPLEX (KINASE/PEPTIDE) (7 CHAIN: A; IIDE HO)YQPQPA; N: B; ILCK	YROSINE COMPLEX (KINASE/PEPTIDE) (7 CHAIN: A; TIDE HO)YQPQPA; N: B; ILCK	YROSINE COMPLEX (TYROSINE
SeqFold Coumpound	GROWTH FACTOR BOUND PROTEIN 2; 1GRI 5 CHAIN: A, B; 1GRI 6	PHOSPHORIC DIESTER HYDROLASE PHOSPHOLIPASE C- GAMMA (SH3 DOMAIN) (E.C.3.1.4.11) IHSQ 3 (NMR, MINIMIZED MEAN STRUCTURE) 1HSQ 4	136.33 P56==LCK== TYROSINE KINASE; ILCK 7 CHAIN: A; ILCK 8 TAIL PHOSPHOPEPTIDE TEGQ(PHOSPHO)YQPQPA; ILCK 14 CHAIN: B; ILCK 15	P56==LCK== TYROSINE KINASE; ILCK 7 CHAIN: A; ILCK 8 TAIL PHOSPHOPEPTIDE TEGQ(PHOSPHO)YQPQPA; ILCK 14 CHAIN: B; ILCK 15	P56==LCK== TYROSINE KINASE; ILCK 7 CHAIN: A; ILCK 8 TAIL PHOSPHOPEPTIDE TEGQ(PHOSPHO)YQPQPA; ILCK 14 CHAIN: B; ILCK 15	95.11 HUMAN P56 TYROSINE KINA SE-11 KK 7 CHAIN- A-
PMF S	0.92	-0.13	<u> </u>	1.00	00.1	96
Verify Score S	0.41	0.04		0.55	0.66	
PSI BLAST Score	1.5e-23 (5.16-10	1.4e-35	6.8e-35	1.4e-35	5.1e-24
End	180	76	193	161	177	193
Start AA	36	30	34	35	38	8
Chain ID	A		٧	A	¥	A
PDB ID	lgri	lhsq	llck	lick	llck	11K
SEQ ID NO:	448	448	448	448	44 8	448

SEQ ID	PDB ID	Chain ID	Start AA	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation	
					Score				PHOSPHOTYROSYL PEPTIDE AC-PTYR-GLU- GLU-ILE; ILKK 11 CHAIN: B: ILKK 12		
448	1lkk	V		191	5.1e-24	1.31	1.00		HUMAN P56 TYROSINE KINASE; 1LKK 7 CHAIN: A; 1LKK 8 PHOSPHOTYROSYL PEPTIDE AC-PTYR-GLU- GLU-ILE; 1LKK 11 CHAIN: B: 1LKK 12	COMPLEX (TYROSINE KINASE/PEPTIDE)	
448	lqcf	∢	37	207	1.2e-44	0.74	1.00		HAEMATOPOETIC CELL KINASE (HCK); CHAIN: A;	TYROSINE KINASE TYROSINE KINASE-INHIBITOR COMPLEX, DOWN-REGULATED KINASE, 2 ORDERED ACTIVATION LOOP	
448	lsem	∢	35	98	1.7e-10	0.09	0.01		SEM-5; ISEM 3 CHAIN: A, B; ISEM 5 10-RESIDUE PROLINE-RICH PEPTIDE FROM MSOS ISEM 8 CHAIN: C, D ISEM 10	SIGNAL TRANSDUCTION PROTEIN SRC-HOMOLOGY 3 (SH3) DOMAIN, PEPTIDE-BINDING PROTEIN, ISEM 18 2 GUANINE NUCLEOTIDE EXCHANGE FACTOR ISEM 19	
448	Isha	Α .	. 16	193	6.8e-25			84.13	PHOSPHOTRANSFERASE V-SRC TYROSINE KINASE TRANSFORMING PROTEIN (PHOSPHOTYROSINE ISHA 3 RECOGNITION DOMAIN SH2) (E.C.2.7.1.112) COMPLEX WITH 1SHA 4 PHOSPHOPEPTIDE A (TYR- VAL-PRO-MET-LEU, PHOSPHORYLATED TYR) ISHA 5		
448	lsha	A	93	192	6.8e-25	1.14	1.00		PHOSPHOTRANSFERASE V-SRC TYROSINE KINASE		

d PDB annotation	PROTEIN NE TON PLEX A (TYR- U,	NASE; TRANSFERASE TRANSFERASE, TYROSINE KINASE, SH3, SH2, ONCOPROTEIN		NULL; TRANSFERASE HCK, SH2, TYROSINE KINASE, SIGNAL TRANSDUCTION, TRANSFERASE		ORT TRANSPORT PP15, B2; TRANSPORT,		DULIN- KINASE KINASE, SIGNAL EIN TRANSDUCTION, ULL; CALCIUM/CALMODULIN	
Coumpound	TRANSFORMING PROTEIN (PHOSPHOTYROSINE 1SHA 3 RECOGNITION DOMAIN SH2) (E.C.2.7.1.112) COMPLEX WITH 1SHA 4 PHOSPHOPEPTIDE A (TYR-VAL-PRO-MET-LEU, PHOSPHORYLATED TYR) 1SHA 5	ABL TYROSINE KINASE; CHAIN: NULL;	ABL TYROSINE KINASE; CHAIN: NULL;	HCK SH2; CHAIN: NULL;	HCK SH2; CHAIN: NULL;	NUCLEAR TRANSPORT	NUCLEAR TRANSPORT FACTOR-2; CHAIN: A, B;	CALCIUM/CALMODULIN- DEPENDENT PROTEIN KINASE; CHAIN: NULL;	U2 RNA HAIRPIN IV; CHAIN: Q, R; U2 A';
SeqFold Score		87.30			103.54	60.79		19:69	
PMF Score			0.99	1.00			96.0		1.00
Verify Score			0.74	0.95			0.49		0.42
PSI BLAST Score		3.4e-29	3.4e-29	3.4e-26	3.4e-26	2.2e-31	2.2e-31	5.4e-25	8.1e-09
End AA		189	192	195	195	141	136	347	397
Start AA		28	59	68	06	12	18	29	341
Chain ID						А	A		В
PDB ID		2abl	2abi	3hck	3hck	lar0	1ar0	1a06	la9n
SEQ ID NO:		448	448	448	448	454	454	457	457

PDB annotation		PROTEIN KINASE CDK2; PROTEIN	KINASE, CELL CYCLE, PHOSPHORYI ATION	STAUROSPORINE, 2 CELL DIVISION.	MITOSIS, INHIBITION	PROTEIN KINASE CDK2; PROTEIN	KINASE, CELL CYCLE,	PHOSPHORYLATION,	STAUROSPORINE, 2 CELL DIVISION, MITOSIS, INHIBITION	COMPLEX (KINASE/INHIBITOR)	CDK6; P19INK4D; CYCLIN	DEPENDENT KINASE, CYCLIN	DEPENDENT KINASE INHIBITORY 2	PROTEIN, CDK, INK4, CELL CYCLE,	COMPLEX (KINASE/INHIBITOR)	HEADER HELIX	COMPLEX (KINASE/INHIBITOR)	CDK6; P19INK4D; CYCLIN	DEPENDENT KINASE, CYCLIN	DEPENDENT KINASE INHIBITORY 2	PROTEIN, CDK, INK4, CELL CYCLE,	COMPLEX (KINASE/INHIBITOR)	HEADER HELIX	COMPLEX (INHIBITOR	PROTEIN/KINASE) INHIBITOR	PROTEIN, CYCLIN-DEPENDENT	KINASE, CELL CYCLE 2 CONTROL,	ALPHA/BETA, COMPLEX (INHIBITOR	PROTEIN/KINASE)	COMPLEX (INHIBITOR	PROTEIN/KINASE) INHIBITOR	PROTEIN, CYCLIN-DEPENDENT
Coumpound	CHAIN: B, D;	CYCLIN-DEPENDENT	PROTEIN KINASE 2;	ישטאוי ואסדיי		CYCLIN-DEPENDENT	PROTEIN KINASE 2;	CHAIN: NULL;		CYCLIN-DEPENDENT	KINASE 6; CHAIN: A, C;	CYCLIN-DEPENDENT	KINASE INHIBITOR;	CHAIN: B, D;			CYCLIN-DEPENDENT	KINASE 6; CHAIN: A, C;	CYCLIN-DEPENDENT	KINASE INHIBITOR;	CHAIN: B, D;			CYCLIN-DEPENDENT	KINASE 6; CHAIN: A;	P19INK4D; CHAIN: B;				CYCLIN-DEPENDENT	KINASE 6; CHAIN: A;	PI9INK4D; CHAIN: B;
SeqFold Score		92.24								86.92									_					103.69		-						_
PMF Score						1.00											0.99													1.00		
Verify Score						0.37											0.42					-								0.32		
PSI BLAST Score		8.1e-33				8.1e-33				5.4e-36							5.4e-36						;	2.2e-39			-		_	2.2e-39		
End AA		317				310				305							304						;	314						304		
Start AA		22				23				21							24						,	91						23		
Chain ID			-							٧							4							Ą						٧		_
PDB ID		laq1				ladi	_			16i8							1bi8							Iblx						1blx		
SEQ NO.		457				457				457							457							457	·					457		

PDB annotation	KINASE, CELL CYCLE 2 CONTROL, ALPHA/BETA, COMPLEX (INHIBITOR PROTEIN/KINASE)	PHOSPHOTRANSFERASE PROTEIN KINASE ICKI 18	PHOSPHOTRANSFERASE PROTEIN KINASE ICKI 18	TRANSFERASE STRESS-ACTIVATED PROTEIN KINASE-3, ERK6, ERK5; P38-GAMMA, GAMMA, PHOSPHORYI ATION MAP KINASE			RNA BINDING PROTEIN/RNA NESTED DOUBLE PSEUDOKNOT RNA STRUCTURE	RNA BINDING PROTEIN RNA- BINDING DOMAIN	PHOSPHOTRANSFERASE FGFRIK, FIBROBLAST GROWTH FACTOR RECEPTOR 1; TRANSFERASE, TYROSINE-PROTEIN KINASE, ATP- BINDING, 2 PHOSPHORYLATION,
Coumpound		CASEIN KINASE I DELTA; ICKI 6 CHAIN: A, B; ICKI 7	CASEIN KINASE I DELTA; ICKI 6 CHAIN: A. B. ICKI 7	PHOSPHORYLATED MAP KINASE P38-GAMMA; CHAIN: A, B;	PHOSPHOTRANSFERASE CAMP-DEPENDENT PROTEIN KINASE CATALYTIC SUBUNIT ICMK 3 (E.C.2.7.1.37)	TRANSFERASE(PHOSPHO TRANSFERASE) CAMP- DEPENDENT PROTEIN KINASE (E.C.2.7.1.37) (CAPK) 1CTP 3 (CATALYTIC SUBUNIT) 1CTP 4	UIA PROTEIN; CHAIN: A; HDV RIBOZYME SELF- CLEAVED; CHAIN: B;	HU ANTIGEN C; CHAIN: A;	FGF RECEPTOR 1; CHAIN: A, B;
SeqFold Score		63.66			84.57	83.76			79.13
PMF Score			0.93	1.00			0.71	0.93	
Verify Score			0.30	0.54			0.26	0.28	
PSI BLAST Score		5.4e-30	5.4e-30	2.4e-36	1.9e-28	8.1e-28	5.4e-09	5.4e-07	2.7e-20
End		324	310	314	357	357	397	397	311
Start AA		16	19	17	e	_	336	345	6
Chain ID		А	٧	∀	ы	ш	A	Ą	¥
PDB ID		1cki	1cki	lcm8	lcmk	lctp	1cx0	1d8z	lfgk
SEQ ID NO:		457	457	457	457	457	457	457	457

SEQ ID NO:	PDB ID	Chain ID	Start	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
┅										RECEPTOR, PHOSPHOTRANSFERASE
	lfgk	B	9	310	1.4e-21			81.51	FGF RECEPTOR 1; CHAIN: A, B;	PHOSPHOTRANSFERASE FGFR1K, FIBROBLAST GROWTH FACTOR RECEPTOR 1; TRANSFERASE,
										TYROSINE-PROTEIN KINASE, ATP- BINDING, 2 PHOSPHORYLATION, RECEPTOR, PHOSPHOTRANSFERASE
1	⊒		341	397	1.4e-08	0.50	0.75		UI SMALL NUCLEAR RIBONUCLEOPROTEIN A; CHAIN: NIII.:	RIBONUCLEOPROTEIN UIA117; RIBONUCLEOPROTEIN, RNP DOMAIN, SPI ICEOSOME
1	of J	4	345	412	8.1e-07	-0.12	0.16		NUCLEOLIN RBD2; CHAIN: A;	STRUCTURAL PROTEIN PROTEIN C23; RNP, RBD, RRM, RNA BINDING POMARIA MICT FOR ITS
	lhcl		22	317	8.1e-35			99.28	HUMAN CYCLIN- DEPENDENT KINASE 2; CHAIN: NULL;	PROTEIN KINASE CDK2; TRANSFERASE, SERINE/THREONINE PROTEIN KINASE, ATP-BINDING, 2 CELL CYCLE, CELL DIVISION, MITOSIS PHOSPHORYI ATION
·	lhcl		23	304	8.1e-35	0.40	1.00		HUMAN CYCLIN- DEPENDENT KINASE 2; CHAIN: NULL;	PROTEIN KINASE CDK2; TRANSFERASE, SERINE/THREONINE PROTEIN KINASE, ATP-BINDING, 2 CELL CYCLE, CELL DIVISION, MITOSIS, PHOSPHORYLATION
}	lian		1	350	8.1e-25			97.38	P38 MAP KINASE; CHAIN: NULL;	SERINE/THREONINE-PROTEIN KINASE CSBP, RK, P38; PROTEIN SER/THR-KINASE, SER/THREONINE-PROTEIN KINASE
	lir3	٧	6	324	2.7e-19			74.34	INSULIN RECEPTOR; CHAIN: A; PEPTIDE SUBSTRATE; CHAIN: B;	COMPLEX (TRANSFERASE/SUBSTRATE) TYROSINE KINASE, SIGNAL TRANSDUCTION, PHOSPHOTRANSFERASE, 2 COMPLEX (KINASE/PEPTIDE SUBSTRATE/ATP

PDB ID	Chain ID	Start AA	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
				Score					ANALOG), ENZYME, 3 COMPLEX
1		12	310	1 1e-31	0.44	200		C-IIIN N-TERMINAI	TRANSFERASE/SUBSTRATE)
		:	3		;	}		KINASE; CHAIN: NULL;	INK3 MAP KINASE,
									SERINE/THREONINE PROTEIN 2 KINASE
1 jnk		4	345	1.1e-31			95.25	C-JUN N-TERMINAL	TRANSFERASE JNK3; TRANSFERASE,
								KINASE; CHAIN: NULL;	JNK3 MAP KINASE, SERINE/THREONINE PROTEIN 2 KINASE
1koa			417	8.1e-29			78.60	TWITCHIN; CHAIN: NULL;	KINASE KINASE, TWITCHIN, INTRASTERIC REGIL ATION
lpme		22	321	1.1e-33	0.32	0.99		ERK2; CHAIN: NULL;	TRANSFERASE MAP KINASE,
			_						SERINE/THREONINE PROTEIN KINASE, TRANSFERASE
1pme		8	341	1.1e-33			94.17	ERK2; CHAIN: NULL;	TRANSFERASE MAP KINASE,
									SERINE/THREONINE PROTEIN KINASE, TRANSFERASE
1qm9	Ą	345	397	8.1e-08	0.19	0.62		POLYPYRIMIDINE TRACT-	RIBONUCLEOPROTEIN PTB, PTB-
								BINDING PROTEIN; CHAIN: A:	C198, HETEROGENEOUS NUCLEAR POLYPYRIMIDINE TRACT BINDING
									PROTEIN, RNP, RNA, SPICING, 2
똢	A	17	365	1.4e-29			96.61	TITIN: CHAIN: A. B:	SERINE KINASE SERINE KINASE.
ı									TITIN, MUSCLE, AUTOINHIBITION
1‡;	⋖	74	304	1.4e-29	0.05	86.0		TITIN; CHAIN: A, B;	SERINE KINASE SERINE KINASE,
1									TITIN, MUSCLE, AUTOINHIBITION
In I	∢	336	397	2.7e-09	0.75	0.76		UIA SPLICEOSOMAL PROTEIN; IURN 5 CHAIN:	COMPLEX (RIBONUCLEOPROTEIN/RINA)
								A, B, C; 1URN 6 RNA	
								21MER HAIRPIN (5'-	
								(AP*AP*UP*CP*CP*AP*UP	
				_				*UP* 10KN 11 CHAIN: P, Q, R 11RN 13	
1									

PDB annotation	RNA-BINDING PROTEIN SPLICING, UZ SNRNP, RBD, RNA-BINDING PROTEIN	TRANSFERASE MITOGEN ACTIVATED PROTEIN KINASE, MAP 2, ERK2; TRANSFERASE, SERINE/THREONINE-PROTEIN KINASE, MAP KINASE, 2 ERK2	TRANSFERASE MITOGEN ACTIVATED PROTEIN KINASE, MAP 2, ERK2; TRANSFERASE, SERINE/THREONINE-PROTEIN KINASE, MAP KINASE, 2 ERK2	GENE REGULATION WINGED HELIX, DNA-RECOGNITION HELIX	DNA BINDING DOMAIN DNA BINDING DOMAIN, WINGED HELIX	GENE REGULATION/DNA HEPATOCYTE NUCLEAR FACTOR 3 FORKHEAD HOMOLOG 2, NMR, STRUCTURE, DYANAMICS, GENESIS, WINGED HELIX PROTEIN, 2 GENE REGULATION/DNA	GENE REGULATION/DNA HEPATOCYTE NUCLEAR FACTOR 3 FORKHEAD HOMOLOG 2, NMR, STRUCTURE, DYANAMICS, GENESIS, WINGED HELIX PROTEIN, 2 GENE REGULATION/DNA	HNF-3 HOMOLOGUES HFH-2; HNF-3 HOMOLOGUES, WINGED HELIX PROTEIN
Coumpound	SPLICING FACTOR UZAF 65 KD SUBUNIT; CHAIN: A;	EXTRACELLULAR REGULATED KINASE 2; CHAIN: NULL;	EXTRACELLULAR REGULATED KINASE 2; CHAIN: NULL;	S12 TRANSCRIPTION FACTOR (FKH-14); CHAIN: A;	AFX; CHAIN: A;	HNF3/FH TRANSCRIPTION FACTOR GENESIS; CHAIN: A; 5'- CHAIN: B; 5'- CHAIN: C;	HNF3/FH TRANSCRIPTION FACTOR GENESIS; CHAIN: A; 5'- CHAIN: B; 5'- CHAIN: C;	GENESIS; CHAIN: NULL;
SeqFold Score		94.87					79.14	
PMF Score	0.78		1.00	00.1	1.00	1.00		1.00
Verify Score	0.18		0.39	0.29	0.43	0.27		0.17
PSI BLAST Score	8.Īe-07	5.4e-34	5.4e-34	8.1e-28	8.1e-26	2.2e-28	2.2e-28	1.6e-28
End AA	397	349	321	179	173	173	197	173
Start AA	345	01	24	100	101	100	100	100
Chain ID	<			٧	¥	Ą	4	
PDB ID	2u2f	3erk	3erk	1d5v	le17	2hdc	2hdc	2hfh
SEQ NO:	457	457	457	458	458	458	458	458

SEQ BOS BOS BOS BOS BOS BOS BOS BOS BOS BOS	PDB CI	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
458	2hfh		66	192	1.6e-28		_	76.10	GENESIS; CHAIN: NULL;	HNF-3 HOMOLOGUES HFH-2; HNF-3 HOMOLOGUES, WINGED HELIX PROTEIN
460	lor5	4	17	118	0.0054	-0.29	0.03		SEC18P (RESIDUES 22 - 210); CHAIN: A, B, C;	ENDOCYTOSIS/EXOCYTOSIS DOUBLE-PSI BETA BARREL, VESICLE FUSION, 2 ENDOCYTOSIS/EXOCYTOSIS
460	lpcf	∀	62	128	5.4e-20			81.97	TRANSCRIPTIONAL COACTIVATOR PC4; CHAIN: A, B, C, D, E, F, G, H;	TRANSCRIPTION P15; TRANSCRIPTION, TRANSCRIPTIONAL COFACTOR, TRANSCRIPTIONAL 2 CO- ACTIVATOR, SSDNA BINDING, NUCLEAR PROTEIN
460	lpcf	V	63	110	5.4e-20	0.33	1.00		TRANSCRIPTIONAL COACTIVATOR PC4; CHAIN: A, B, C, D, E, F, G, H;	TRANSCRIPTION P15; TRANSCRIPTION, TRANSCRIPTIONAL COFACTOR, TRANSCRIPTIONAL 2 CO- ACTIVATOR, SSDNA BINDING, NUCLEAR PROTEIN
461	lcmz	∢	75	202	2.7e-50			110.58	GAIP (G-ALPHA INTERACTING) PROTEIN; CHAIN: A;	SIGNALING PROTEIN REGULATION GALPHA INTERACTING PROTEIN; GAIP, RGS, REGULATOR OF G PROTEIN, SIGNALING PROTEIN 2 REGULATION
461	lcmz	¥	9/	202	2.7e-50	0.54	00.1		GAIP (G-ALPHA INTERACTING) PROTEIN; CHAIN: A;	SIGNALING PROTEIN REGULATION GALPHA INTERACTING PROTEIN; GAIP, RGS, REGULATOR OF G PROTEIN, SIGNALING PROTEIN 2 REGULATION
461	1dk8	¥	78	209	5.4e-42	0.43	0.98		AXIN; CHAIN: A;	SIGNALING PROTEIN ALPHA-HELIX, PI-HELIX
461	lemu	A	82	200	1.6e-37	0.50	1.00		AXIN, CHAIN: A; ADENOMATOUS POLYPOSIS COLI	SIGNALING PROTEIN RGS DOMAIN

PDB annotation		TOXIN CYTOXIN (CARDIOTOXIN), MEMBRANE PERTURBATION, CIS/TRANS 2 ISOMERIZATION, BOUND WATER					OXIDOREDUCTASE FERROCYTOCHROME C'OXYGEN OXIDOREDUCTASE; OXIDOREDUCTASE, CYTOCHROME(C)-OXYGEN, CYTOCHROME C 2 OXIDASE	OXIDOREDUCTASE FERROCYTOCHROME C\:\text{OXYGEN} OXIDOREDUCTASE; OXIDOREDUCTASE, CYTOCHROME(C)-OXYGEN, CYTOCHROME C 2 OXIDASE
Coumpound	PROTEIN; CHAIN: B;	CYTOTOXIN 2; CHAIN: A	CYTOTOXIN CARDIOTOXIN V=4==_/II\$= (TOXIN /III\$) ICDT 3	CYTOTOXIN TOXIN GAMMA (CARDIOTOXIN) ITGX 3	POSTSYNAPTIC NEUROTOXIN ALPHA- *BUNGAROTOXIN 2ABX 4	CARDIOTOXIN CARDIOTOXIN CTX I (NMR, 11 STRUCTURES) 2CDX 3	CYTOCHROME C OXIDASE; CHAIN: A, B, C, D, E, F, G, H, I, J, K, L, M, N, O, P, Q,	CYTOCHROME C OXIDASE; CHAIN: A, B, C, D, E, F, G, H, I, J, K, L, M, N, O, P, Q,
SeqFold Score							57.62	
PMF Score		0.35	0.00	0.09	0.05	0.01		0.34
Verify Score		-0.44	0.23	-0.11	-0.03	0.00		-0.89
PSI BLAST Score		0.0081	0.0054	0.0054	0.00081	0.0054	2.7e-20	2.7e-20
End AA		99	99	99	171	99	120	11.8
Start AA		\$	5	\$	16	5	64	73
Chain ID		V	∢	A	A		T.	-
PDB ID		1cb9	1cdt	ltgx	2арх	2cdx	20cc	20cc
SEQ ID NO:		463	463	463	463	463	465	465

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PDB annotation	COMPLEX (NUCLEAR PROTEIN/RNA), COMPLEX (NUCLEAR PROTEIN/RNA), RNA, SNRNP,RIBONUCLEOPROTEIN	COMPLEX (NUCLEAR PROTEIN/RNA), COMPLEX (NUCLEAR PROTEIN/RNA), RNA, SNRNP, RIBONUCLEOPROTEIN	RNA-BINDING PROTEINRNA TRA PRE-MRNA; SPLICING REGULATION, RNP DOMAIN, RNA COMPLEX	RNA-BINDING PROTEINRNA TRA PRE-MRNA, SPLICING REGULATION, RNP DOMAIN, RNA COMPLEX	GENE REGULATIONRNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM,
Coumpound	U2 RNA HAIRPIN IV; CHAIN: Q, R; U2 A; CHAIN: A, C; U2 B"; CHAIN: B, D;	U2 RNA HAIRPIN IV; CHAIN: Q, R; U2 A'; CHAIN: A, C; U2 B"; CHAIN: B, D;	SXL-LETHAL PROTEIN; CHAIN: A, B; RNA (5'- R(P*GP*UP*UP*GP*UP*UP *UP*UP*UP*UP*UP*U)- CHAIN: P, Q;	SXL-LETHAL PROTEIN; CHAIN: A, B; RNA (5* R(P*GP*UP*UP*GP*UP*UP *UP*UP*UP*UP*UP*UP*U)- CHAIN: P, Q;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP*AP* AP*AP*AP*AP*A)-3'); CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A)-3); CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN I; CHAIN: A, B,
SeqFold Score		147.88	59.62		54.28		
PMF Score	1.00			0.92		0.86	0.52
Verify Score	1.50			0.18		0.42	0.26
PSI BLAST Score	8.1e-32	8.1e-32	8.1e-16	8.1e-16	1.4e-14	1.4e-14	8.1e-15
End	96	96	172	161	174	173	161
Start AA	۴	٣	S	9	7	∞	8
Chain ID	В	B	٧	∢	A	¥	В
PDB ID	la9n	1a9n	1b7f	1b7f	lcvj	lcvj	lcvj
SEQ ID NO:	467	467	467	467	467	467	467

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PDB annotation	PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	NUCLEAR PROTEIN HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1, NUCLEAR PROTEIN, HNRNP, RBD, RRM, RNP, RNA BINDING, 2 RIBONUCLEOPROTEIN	RIBONUCLEOPROTEIN PTB, PTB. C198, HETEROGENEOUS NUCLEAR POLYPYRIMIDINE TRACT BINDING PROTEIN, RNP, RNA, SPICING, 2 TRANSLATION	NUCLEAR PROTEIN UI SNRNP A PROTEIN; RNA BINDING DOMAIN, NICLEAR PROTEIN	NUCLEAR PROTEIN UI SNRNP A
Coumpound	C, D, E, F, G, H; RNA (5- R(*AP*AP*AP*AP*AP*AP* AP*AP*AP*AP*A)-3'); CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP3); CHAIN: M, N, O, P, Q, R, S, T;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*A)-3); CHAIN: M, N, O, P, Q, R, S, T;	HNRNP A1; CHAIN: NULL;	POLYPYRIMIDINE TRACT- BINDING PROTEIN; CHAIN: A;	UI SMALL NUCLEAR RIBONUCLEOPROTEIN A; CHAIN: NULL:	UI SMALL NUCLEAR
SeqFold Score							64.11
PMF Score		0.99	0.89	0.78	0.33	0.99	
Verify Score		0.26	0.47	0.41	-0.21	-0.11	
PSI BLAST Score		1.9e-12	8.1e-14	1.1e-10	8.1e-21	1.9e-09	1.9e-09
End		149	151	167	174	174	174
Start AA		∞	8	∞	9	127	25
Chain ID		tr.	Н		A		
PDB ID		lcvj	Icvj	lhal	lqm9	2u la	2ula
SEQ ID NO:		467	467	467	467	467	467

nd PDB annotation	OTEIN A; PROTEIN; RNA BINDING DOMAIN, NUCLEAR PROTEIN	COMPLEX (RIBONUCLEOPROTEIN/DNA) HNRNP OTEIN A1, UP1; COMPLEX (RIBONUCLEOPROTEIN/DNA), HETEROGENEOUS NUCLEAR 2 DMETRIC RIBONUCLEOPROTEIN A1	HATASE SCAFFOLD PROTEIN SCAFFOLD B; PROTEIN, PP2A, PHOSPHORYLATION, HEAT REPEAT	ANT SUGAR BINDING PROTEIN C-TYPE JN: A, B, LECTIN, CRD, SP-D, COLECTIN, ALPHA-HELICAL COILED- 2 COIL, LUNG SURFACTANT, SUGAR BINDING PROTEIN		JLL; NK CELL NK CELL, RECEPTOR, C- TYPE LECTIN, C-TYPE LECTIN-LIKE, NKD	ALPHA MEMBRANE PROTEIN C-TYPE I. A; LECTIN-LIKE DOMAINS BETA I. B	TON HEMATOPOIETIC CELL RECEPTOR CHAIN: A; ACTIVATION INDUCER MOLECULE (AIM), EA 1, HEMATOPOIETIC CELL RECEPTOR, LEUCOCYTE, C-TYPE
Coumpound	RIBONUCLEOPROTEIN A; CHAIN: NULL;	HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1; CHAIN: A; 12- NUCLEOTIDE SINGLE- STRANDED TELOMETRIC DNA; CHAIN: B;	PROTEIN PHOSPHATASE PP2A; CHAIN: A, B;	LUNG SURFACTANT PROTEIN D; CHAIN: A, B, C;	CD94; CHAIN: NULL;	CD94; CHAIN: NULL;	FLAVOCETIN-A: ALPHA SUBUNIT; CHAIN: A; FLAVOCETIN-A: BETA SUBUNIT; CHAIN: B	EARLY ACTIVATION ANTIGEN CD69; CHAIN: A;
SeqFold Score						86.24		
PMF Score		0.24	0.15	0.19	1.00		0.78	0.87
Verify Score		0.28	-0.15	0.02	0.53		90.0	0.55
PSI BLAST Score		2.4e-12	0.0011	1.1e-23	5.4e-26	5.4e-26	2.7e-24	8.1e-26
End AA		174	444	258	260	261	261	259
Start AA		∞	222	100	134	134	137	135
Chain ID		∢	A	∢			æ	¥
PDB ID		2up1	1b3u	1608	1566	166e	1c3a	1e87
SEQ ID NO:		467	468	469	469	469	469	469

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PDB annotation	COAGULATION FACTOR BINDING IXX-BP COAGULATION FACTOR BINDING, C-TYPE LECTIN, GLA- DOMAIN 2 BINDING, C-TYPE CRD MOTIF, LOOP EXCHANGED DIMFR	COAGULATION FACTOR BINDING IXX-BP COAGULATION FACTOR BINDING, C-TYPE LECTIN, GLA- DOMAIN 2 BINDING, C-TYPE CRD MOTE 1,00P EXCHANGED DIMER	PANCREATIC STONE INHIBITOR PANCREATIC STONE INHIBITOR, LECTIN	METAL BINDING PROTEIN PANCREATIC STONE PROTEIN, PSP; PANCREATIC STONE INHIBITOR, LITHOSTATHINE	METAL BINDING PROTEIN PANCREATIC STONE PROTEIN, PSP; PANCREATIC STONE INHIBITOR, LITHOSTATHINE	COMPLEX (NK RECEPTOR/MHC CLASS I) H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, B2M; NK-CELL SURFACE GLYCOPROTEIN YEI/48, NK CELL, INHIBITORY RECEPTOR, MHC-I, C- TYPE LECTIN-LIKE, 2 HISTOCOMPATIBILITY, B2M, LY49, LY-49	COMPLEX (NK RECEPTOR/MHC CLASS I) H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, B2M; NK-CELL SURFACE
Coumpound	COAGULATION FACTORS IX/X-BINDING PROTEIN; CHAIN: A, B, C, D, E, F;	COAGULATION FACTORS IX/X-BINDING PROTEIN; CHAIN: A, B, C, D, E, F;	LITHOSTATHINE; CHAIN: NULL	LITHOSTATHINE; CHAIN: A;	LITHOSTATHINE; CHAIN: A;	MHC CLASS I H-2DD HEAVY CHAIN; CHAIN: A; BETA-2-MICROGLOBULIN; CHAIN: B; HIV ENVELOPE GLYCOPROTEIN 120 PEPTIDE; CHAIN: P; LY49A; CHAIN: C, D;	MHC CLASS I H-2DD HEAVY CHAIN; CHAIN: A; BETA-2-MICROGLOBULIN; CHAIN: B; HIV ENVELOPE
SeqFold Score	56.80		53.00	59.68			
PMF Score		0.62			96.0	0.75	1.00
Verify Score		0.18			0.36	0.24	0.14
PSI BLAST Score	1.3e-23	1.3e-23	1.9e-21	1.4e-24	1.4c-24	1.4e-27	2.7e-25
End AA	261	261	261	261	260	260	260
Start AA	136	137	137	124	134	132	141
Chain ID	В	В		A	A	ပ	Q
PDB ID	Lix X	lixx	Uit	1qdd	1qdd	1403	1403
SEQ NO:	469	469	469	469	469	469	469

PDB annotation	GLYCOPROTEIN YE1/48, NK CELL, INHIBITORY RECEPTOR, MHC-1, C- TYPE LECTIN-LIKE, 2 HISTOCOMPATIBILITY, B2M, LY49, LY-49	LECTIN TETRANECTIN, PLASMINOGEN BINDING, KRINGLE 4, C-TYPE LECTIN, 2 CARBOHYDRATE RECOGNITION DOMAIN	ANTIFREEZE PROTEIN RECOMBINANT SEA RAVEN PROTEIN, SOLUTION BACKBONE FOLD, C- 2 TYPE LECTIN, ANTIFREEZE PROTEIN		NK CELL NK CELL, RECEPTOR, C- TYPE LECTIN, C-TYPE LECTIN-LIKE, NKD	NK CELL NK CELL, RECEPTOR, C- TYPE LECTIN, C-TYPE LECTIN-LIKE, NKD	MEMBRANE PROTEIN C-TYPE LECTIN-LIKE DOMAINS	A; ACTIVATION INDUCER MOLECULE (AIM), EA 1, HEMATOPOIETIC CELL RECEPTOR, LEUCOCYTE, C-TYPE LECTIN-LIKE, 2 NKD, KLR
Coumpound	GLYCOPROTEIN 120 PEPTIDE; CHAIN: P; LY49A; CHAIN: C, D;	TETRANECTIN; CHAIN: NULL;	SEA RAVEN TYPE II ANTIFREEZE PROTEIN; CHAIN: A;	LUNG SURFACTANT PROTEIN D; CHAIN: A, B, C;	CD94; CHAIN: NULL;	CD94; CHAIN: NULL;	FLAVOCETIN-A: ALPHA SUBUNIT; CHAIN: A; FLAVOCETIN-A: BETA SUBUNIT; CHAIN: B	EARLY ACTIVATION ANTIGEN CD69; CHAIN: A;
SeqFold Score						86.29		
PMF		0.89	0.58	0.25	1.00		0.78	0.87
Verify Score		0.15	-0.06	-0.05	0.53		90.0	0.55
PSI BLAST Score		2.7e-24	5.4e-26	5.4e-24	5.4e-26	5.4e-26	2.7e-24	8.1e-26
End			258	285	287	288	288	286
Start AA		134	132		161	191	164	162
Chain ID			Ą	A			В	A
PDB ED		1tn3	2afp	1608	1b6e	1666	1c3a	1e87
SEQ NO:		469	469	469	469	469	469	469

PDB annotation	COAGULATION FACTOR BINDING IXX-BP COAGULATION FACTOR BINDING, C-TYPE LECTIN, GLA- DOMAIN 2 BINDING, C-TYPE CRD MOTIF. LOOP EXCHANGED DIMER	COAGULATION FACTOR BINDING IXX-BP COAGULATION FACTOR BINDING, C-TYPE LECTIN, GLA- DOMAIN 2 BINDING, C-TYPE CRD MOTIF, LOOP EXCHANGED DIMER	COAGULATION FACTOR BINDING IXX-BP COAGULATION FACTOR BINDING, C-TYPE LECTIN, GLA- DOMAIN 2 BINDING, C-TYPE CRD MOTIF, LOOP EXCHANGED DIMER	PANCREATIC STONE INHIBITOR, PANCREATIC STONE INHIBITOR, LECTIN	METAL BINDING PROTEIN PANCREATIC STONE PROTEIN, PSP; PANCREATIC STONE INHIBITOR, LITHOSTATHINE	METAL BINDING PROTEIN PANCREATIC STONE PROTEIN, PSP; PANCREATIC STONE INHIBITOR,	COMPLEX (NK RECEPTOR/MHC CLASS I) H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, B2M; NK-CELL, SURFACE GLYCOPROTEIN YE1/48, NK CELL, INHIBITORY RECEPTOR, MHC-1, C- TYPE LECTIN-LIKE, 2 HISTOCOMPATIBILITY, B2M, LY49,
Coumpound	CHAIN: A, B, C, D, E, F; D CHAIN: A, B, C, D, E, F; D M	COAGULATION FACTORS C IX/X-BINDING PROTEIN; D CHAIN: A, B, C, D, E, F; B D M	COAGULATION FACTORS IXX-BINDING PROTEIN; CHAIN: A, B, C, D, E, F; D M	LITHOSTATHINE; CHAIN: P. NULL LI	LITHOSTATHINE; CHAIN: P, A; P, L,	LITHOSTATHINE; CHAIN: PA A; PA LI	MHC CLASS I H-2DD GHEAVY CHAIN; CHAIN: A; CI BETA-2-MICROGLOBULIN; H CHAIN: B; HIV ENVELOPE GLYCOPROTEIN 120 GPEPTIDE; CHAIN: P; I Y49A; CHAIN: C, D; H
SeqFold Score	50.25	58.01		54.16	63.08		
PMF Score			0.62			96.0	0.72
Verify Score			0.18			0.36	0.41
PSI BLAST Score	8.1e-21	1.3e-23	1.3e-23	1.9e-21	1.4e-24	1.4e-24	8.16-28
End AA	286	288	288	288	288	287	287
Start AA	163	163	164	164	151	191	157
Chain ID	A	В	В		V	A	ပ
PDB ID	lixx	lixx	lixx	11it	1qdd	1qdd	1403
SEQ NO:	469	469	469	469	469	469	469

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PDB annotation	LY-49	COMPLEX (NR RECEPTOR/MHC CLASS I) H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN.	B2M; NK-CELL SURFACE	GLYCOPROTEIN YE1/48, NK CELL, INHIBITORY RECEPTOR, MHC-1. C-	TYPE LECTIN-LIKE, 2 HISTOCOMPATIBILITY, B2M, LY49,	LI-49	PLASMINOGEN BINDING, KRINGLE 4, C-TYPE LECTIN, 2 CARBOHYDRATE RECOGNITION DOMAIN	ANTIFREEZE PROTEIN	RECOMBINANT SEA RAVEN	PROTEIN, SOLUTION BACKBONE FOI D. C. 2 TYPE I ECTIN	ANTIFREEZE PROTEIN	SUGAR BINDING PROTEIN C-TYPE	LECTIN, CRD, SP-D, COLECTIN,	ALPHA-HELICAL COILED- 2 COIL,	LUNG SUKFACIANI, SUGAK BINDING PROTEIN	NK CELL NK CELL, RECEPTOR, C-	TYPE LECTIN, C-TYPE LECTIN-LIKE, NKD	NK CELL NK CELL, RECEPTOR, C-	TYPE LECTIN, C-1 YPE LECTIN-LIKE, NKD	MEMBRANE PROTEIN C-TYPE	LECTIN-LIKE DOMAINS
Coumpound		MHC CLASS I H-2DD HEAVY CHAIN; CHAIN: A; BETA-2-MICROGLOBULIN:	CHAIN: B; HIV ENVELOPE	GLYCOPROTEIN 120 PEPTIDE: CHAIN: P:	LY49A; CHAIN: C, D;	TETRANECTIN: CHAIN:	NULL;	SEA RAVEN TYPE II	ANTIFREEZE PROTEIN;	CHAIN: A;		LUNG SURFACTANT	PROTEIN D; CHAIN: A, B,	హ		CD94; CHAIN: NULL;		CD94; CHAIN: NULL;		FLAVOCETIN-A: ALPHA	SUBUNIT; CHAIN: A; FLAVOCETIN-A: BETA
SeqFold Score																		86.24			
PMF Score		1.00				0.80		0.58				0.19				1.00				0.78	
Verify Score		0.14				0.15		-0.06				0.02				0.53				90.0	
PSI BLAST Score		2.7e-25				2 7F-24		5.4e-26				1.1e-23	-			5.4e-26		5.4e-26		2.7e-24	
End		287				285	}	285				258				260		261		261	
Start AA		891				191		159				001				134		134		137	
Chain ID		Q						¥.				A								В	
PDB ID		1903				1tn3		2afp				1608				1b6e		1b6e		lc3a	
SEQ ID NO:		469				469		469				470				470		470		470	

PDB annotation		HEMATOPOIETIC CELL RECEPTOR ACTIVATION INDUCER MOLECULE (AIM), EA 1, HEMATOPOIETIC CELL RECEPTOR, LEUCOCYTE, C-TYPE LECTIV-LIKE 2 NKD KLR			PANCREATIC STONE INHIBITOR PANCREATIC STONE INHIBITOR, LECTIN	METAL BINDING PROTEIN PANCREATIC STONE PROTEIN, PSP; PANCREATIC STONE INHIBITOR, LITHOSTATHINE	METAL BINDING PROTEIN PANCREATIC STONE PROTEIN, PSP; PANCREATIC STONE INHIBITOR, LITHOSTATHINE	
Coumpound	SUBUNIT: CHAIN: B	EARLY ACTIVATION ANTIGEN CD69; CHAIN: A;	COAGULATION FACTORS IXX-BINDING PROTEIN; CHAIN: A, B, C, D, E, F;	COAGULATION FACTORS IXX-BINDING PROTEIN; CHAIN: A, B, C, D, E, F;	LITHOSTATHINE; CHAIN: NULL	LITHOSTATHINE; CHAIN: A;	LITHOSTATHINE; CHAIN: A;	MHC CLASS I H-2DD HEAVY CHAIN; CHAIN: A; BETA-2-MICROGLOBULIN; CHAIN: B; HIV ENVELOPE GLYCOPROTEIN 120 PEPTIDE; CHAIN: P;
SeqFold Score			56.80		53.00	59.68		
PMF Score		0.87		0.62			96.0	0.75
Verify Score		0.55		0.18			0.36	0.24
PSI BLAST	Score	8.1e-26	1.3e-23	1.3e-23	1.9e-21	1.4e-24	1.4e-24	1.4e-27
End		259	261	261	261	261	260	260
Start		135	136	137	137	124	134	132
Chain ID		4	В	Д		V V	V	U
PDB UD		le87	lix	lixx	Ħ	1qdd	1qdd	1403
SEQ EQ	<u> </u>	470	470	470	470	470	470	470

	T	τ															. –		-		_	
PDB annotation	HISTOCOMPATIBILITY, B2M, LY49, LY-49	COMPLEX (NK RECEPTOR/MHC CLASS I) H-2 CLASS I	HISTOCOMPATIBILITY ANTIGEN, B2M; NK-CELL SURFACE	GLYCOPROTEIN YE1/48, NK CELL,	INHIBITORY RECEPTOR, MHC-1, C- TYPE LECTIN-LIKE, 2	HISTOCOMPATIBILITY, B2M, LY49, LY-49	LECTIN TETRANECTIN,	PLASMINOGEN BINDING, KKINGLE 4, C-TYPE LECTIN, 2 CARBOHYDRATE PECOCNITION POWARI	ANTIFREEZE PROTEIN	RECOMBINANT SEA RAVEN	PROTEIN, SOLUTION BACKBONE	FOLD, C- 2 TYPE LECTIN, ANTHER REZE DROTTEN	SUGAR BINDING PROTEIN C-TYPE	LECTIN, CRD, SP-D, COLECTIN,	ALPHA-HELICAL COILED- 2 COIL,	LUNG SURFACTANT, SUGAR BINDING PROTEIN	NK CELL NK CELL, RECEPTOR, C-	TYPE LECTIN, C-TYPE LECTIN-LIKE,	NK CELL NK CELL, RECEPTOR, C-	TYPE LECTIN, C-TYPE LECTIN-LIKE, NKD	MEMBRANE PROTEIN C-TYPE	LECTIN-LIKE DOMAINS
Coumpound		MHC CLASS I H-2DD HEAVY CHAIN; CHAIN: A;	BETA-2-MICROGLOBULIN; CHAIN: B; HIV ENVELOPE	GLYCOPROTEIN 120	PEPTIDE; CHAIN: P; LY49A; CHAIN: C, D;		TETRANECTIN; CHAIN:	NOLL;	SEA RAVEN TYPE II	ANTIFREEZE PROTEIN;	CHAIN: A;		LUNG SURFACTANT	PROTEIN D; CHAIN: A, B,	ပ်		CD94; CHAIN: NULL;		CD94; CHAIN: NULL;		FLAVOCETIN-A: ALPHA	SUBUNIT; CHAIN: A; FLAVOCETIN-A: BETA
SeqFold Score																			86.29			
PMF Score		1.00					68.0		0.58				0.25				1.00				0.78	-
Verify Score		0.14					0.15		-0.06				-0.05				0.53				90.0	
PSI BLAST Score		2.7e-25					2.7e-24		5.4e-26				5.4e-24				5.4e-26		5.4e-26		2.7e-24	
End		260					258		258				285				287		288		288	
Start AA		141					134		132				148				191		161		164	
Chain ID		Ω							A				A								В	
PDB ID		1403					1tn3		2afp				1508				1b6e		1b6e		1c3a	
SEQ ID NO:		470					470		470				470				470		470		420	

E	_		200	PSI	Verify	PMF	SeaFold	Commodina	DND omnototion
	<u>a</u>	¥¥	AA	BLAST Score	Score	Score	Score		
ıł								SUBUNIT; CHAIN: B	
le87	A	162	286	8.1e-26	0.55	0.87		EARLY ACTIVATION ANTIGEN CD69: CHAIN: A:	HEMATOPOIETIC CELL RECEPTOR ACTIVATION INDICER MOI FOILE
									(AIM), EA 1, HEMATOPOIETIC CELL
									RECEPTOR, LEUCOCYTE, C-TYPE
1ixx	¥	163	286	8.1e-21			50.25	COAGULATION FACTORS	COAGULATION FACTOR BINDING
								IX/X-BINDING PROTEIN;	IX/X-BP COAGULATION FACTOR
								CHAIN: A, B, C, D, E, F;	BINDING, C-TYPE LECTIN, GLA-
									MOTIF 1 OOP EXCHANGED SINGED
lixx	В	163	288	1 30,73			50.03	Carono Carona Haran	MOLIF, LOUF EACHANGED DIMER
	à	3	007	1.35-23			10.86	COAGULATION FACTORS IX/X-BINDING PROTEIN;	COAGULATION FACTOR BINDING IX/X-BP COAGULATION FACTOR
								CHAIN: A, B, C, D, E, F;	BINDING, C-TYPE LECTIN, GLA-
									DOMAIN 2 BINDING, C-TYPE CRD
1	,								MOTIF, LOOP EXCHANGED DIMER
ž	8	164	288	1.3e-23	0.18	0.62		COAGULATION FACTORS	COAGULATION FACTOR BINDING
								IX/X-BINDING PROTEIN;	IX/X-BP COAGULATION FACTOR
								CHAIN: A, B, C, D, E, F;	BINDING, C-TYPE LECTIN, GLA-
									DOMAIN 2 BINDING, C-TYPE CRD
		13.	9	;		1			MOTIF, LOOP EXCHANGED DIMER
1		104	887	1.9e-21			54.16	LITHOSTATHINE; CHAIN:	PANCREATIC STONE INHIBITOR
								ואסקק	FANCKEATIC STONE INHIBITOR, LECTIN
Iqdd	∢	151	288	1.4e-24			63.08	LITHOSTATHINE; CHAIN:	METAL BINDING PROTEIN
					-			A;	PANCREATIC STONE PROTEIN, PSP;
									PANCREATIC STONE INHIBITOR,
,		151	200		1	1			LITHOSTATHINE
n da	<		/87	1.4e-24	0.36	96.0		LITHOSTATHINE; CHAIN:	METAL BINDING PROTEIN
						_		Ą;	PANCREATIC STONE PROTEIN, PSP;
									PANCREATIC STONE INHIBITOR,
1403	ပ	157	287	8.1e-28	0.41	0.72		MHC CLASS I H-2DD	COMPLEX ONK RECEPTOR ACHO
- [\dashv				HEAVY CHAIN; CHAIN: A;	CLASS I) H-2 CLASS I

SEQ ID	PDB TD	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
					2000				BETA-2-MICROGLOBULIN; CHAIN: B; HIV ENVELOPE GLYCOPROTEIN 120 PEPTIDE; CHAIN: P; LY49A; CHAIN: C, D;	HISTOCOMPATIBILITY ANTIGEN, B2M; NK-CELL SURFACE GLYCOPROTEIN YE1/48, NK CELL, INHIBITORY RECEPTOR, MHC-I, C- TYPE LECTIN-LIKE, 2 HISTOCOMPATIBILITY, B2M, LY49,
470	1403	Q	168	287	2.7e-25	0.14	1.00		MHC CLASS I H-2DD HEAVY CHAIN; CHAIN: A; BETA-2-MICROGLOBULIN; CHAIN: B; HIV ENVELOPE GLYCOPROTEIN 120 PEPTIDE; CHAIN: P; LY49A; CHAIN: C, D;	COMPLEX (NK RECEPTOR/MHC CLASS I) H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, B2M; NK-CELL SURFACE GLYCOPROTEIN YE1/48, NK CELL, NHIBITORY RECEPTOR, MHC-1, C-TYPE LECTIN-LIKE, 2 HISTOCOMPATIBILITY, B2M, LY49, 1 Y-49
470	1tn3		161	285	2.7e-24	0.15	0.89		TETRANECTIN; CHAIN: NULL;	LECTIN TETRANECTIN, PLASMINOGEN BINDING, KRINGLE 4, C-TYPE LECTIN, 2 CARBOHYDRATE RECOGNITION DOMAIN
470	2afp	Y	159	285	5.4e-26	-0.06	0.58		SEA RAVEN TYPE II ANTIFREEZE PROTEIN; CHAIN: A;	ANTIFREEZE PROTEIN RECOMBINANT SEA RAVEN PROTEIN, SOLUTION BACKBONE FOLD, C- 2 TYPE LECTIN, ANTIFREEZE PROTEIN
472	1qfc	A	64	328	8.1e-19	-0.09	0.57		PURPLE ACID PHOSPHATASE; CHAIN: A;	HYDROLASE TARTRATE RESISTANT ACID PHOSPHATASE, TRAP; HYDROLASE. METAL PHOSPHATASE
472	Iqhw	4	70	335	5.4e-22	-0.03	0.23		PURPLE ACID PHOSPHATASE; CHAIN: A;	HYDROLASE TARTRATE-RESISTANT ACID PHOSPHATASE; METAL PHOSPHATASE, HYDROLASE
475	1bih	A	20	220	5.4e-09	0.28	0.29		HEMOLIN; CHAIN: A, B;	INSECT IMMUNITY INSECT

SEQ NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation	
										IMMUNITY, LPS-BINDING, HOMOPHILIC ADHESION	
475	lcvs	ပ	99	162	5.4e-08	0.10	-0.09		FIBROBLAST GROWTH	GROWTH FACTOR/GROWTH FACTOR	т.
									FACTOR 2; CHAIN: A, B;	RECEPTOR FGF, FGFR,	
									FACTOR RECEPTOR 1:	IMMUNOGLOBULIN-LIKE, SIGNAL	
									CHAIN: C, D;	GROWTH FACTOR/GROWTH FACTOR	
475	lcvs	6	95	163	5 10 07	2	0.10	T	TIPPODI ACT OPONITION	KECEPTOK	-,-
}	2	١	2	701	3.4e-0/	40.04	07.0		FIBROBLASI GROWIN	GROWTH FACTOR/GROWTH FACTOR	
	_								FACTOR Z; CHAIN: A, B;	RECEPTOR FGF, FGFR,	
									TIDROBLASI GROWIN	IMIMONOGLOBOLIN-LIKE, SIGNAL	
									FACTOR RECEPTOR 1; CHAIN: C. D:	GROWTH FACTOR GROWTH FACTOR	
										RECEPTOR	
475	lepf	∢	22	162	8.1e-07	-0.03	0.40		NEURAL CELL ADHESION MOLECULE; CHAIN: A, B.	CELL ADHESION NCAM; NCAM, IMMUNOGLOBULIN FOLD.	
									C, D;	GLYCOPROTEIN	
475	levt	ပ	57	162	2.2e-07	0.43	0.19		FIBROBLAST GROWTH	GROWTH FACTOR/GROWTH FACTOR	_
									FACTOR 1; CHAIN: A, B;	RECEPTOR FGF1; FGFR1;	
									FIBROBLAST GROWTH	IMMUNOGLOBULIN (IG) LIKE	
									FACTOR RECEPTOR 1;	DOMAINS BELONGING TO THE I-SET	
									CHAIN: C, D;	2 SUBGROUP WITHIN IG-LIKE	
75	3		7,7	6	5	,	1	1		DOMAINS, B-TREFOIL FOLD	
	h711	<		239	5.4e-23	0.46	0.96		HIGH AFFINITY	IMMUNE SYSTEM FC-EPSILON RI-	
									IMMONOGLOBULIN	ALPHA; IMMUNOGLOBULIN FOLD,	
							-		EPSILON RECEPTOR	GLYCOPROTEIN, RECEPTOR, IGE-	
+	١		1		+				CHAIN: A;	BINDING 2 PROTEIN	
4/5	Irba		46	231	1.3e-23	0.30	0.45		HIGH AFFINITY	IMMUNE SYSTEM HIGH AFFINITY	
							•		IMMUNOGLOBULIN	IGE-FC RECEPTOR, FC(EPSILON) IGE-	_
_		_		_					EPSILON RECEPTOR	FC; IMMUNOGLOBULIN FOLD,	
									CHAIN: A; IG EPSILON	GLYCOPROTEIN, RECEPTOR, IGE-	
									CHAIN C REGION; CHAIN:	BINDING 2 PROTEIN, IGE ANTIBODY,	
+		1	1	-+		T			B, D;	IGE-FC	
475	lfcg	V V	46	236	8.1e-29	0.32	0.99		FC RECEPTOR	IMMUNE SYSTEM, MEMBRANE	, —,

Coumpound PDB annotation	FC(GAMMA)RIIA; CHAIN: PROTEIN CD32; FC RECEPTOR, IMMUNOGLOULIN, LEUKOCYTE, CD32	TY IMMUNE SYSTEM RECEPTOR BETA SANDWICH, IMMUNOGLOBULIN- LIKE, RECEPTOR	CHAIN: INHIBITORY RECEPTOR KILLER CELL INHIBITORY RECEPTOR; INHIBITORY RECEPTOR, NATURAL KILLER CELLS, IMMUNOLOGICAL 2 RECEPTORS, IMMUNOGLOBULIN FOLD	CHAIN: INHIBITORY RECEPTOR KILLER CELL INHIBITORY RECEPTOR; INHIBITORY RECEPTOR, NATURAL KILLER CELLS, IMMUNOLOGICAL 2 RECEPTORS, IMMUNOGLOBULIN FOLD		ADHESION MOLECULE-1; D1,2; 1VCA 6 IMMUNOGLOBULIN 1VCA 4 CHAIN: A, B; 1VCA 1VCA 15	ADHESION MOLECULE-1; D1,2; 1VCA 6 IMMUNOGLOBULIN 1VCA 4 CHAIN: A, B; 1VCA 1 SUPERFAMILY, INTEGRIN-BINDING 1VCA 15	CULE-2;
	FC(GAMMA) A;	LOW AFFINITY IMMUNOGLOBULIN GAMMA FC REGION CHAIN: A:	PS8-CL42 KIR; CHAIN: NULL;	PS8-CL42 KIR; CHAIN: NULL;	P58-CL42 KIR; CHAIN: NULL;	HUMAN VAS ADHESION M IVCA 4 CHAI 5	HUMAN VAS ADHESION N 1VCA 4 CHAI 5	INTERCELLULAR ADHESION MOLE
SeqFold Score				146.01		52.80		
PMF		0.54	1.00		1.00		0.07	-0.05
Verify Score		0.41	0.51		0.72		0.12	0.17
PSI BLAST	Score	1.6e-28	1.3e-28	2.7e-68	2.7e-68	1.1e-08	1.1e-08	5.4e-08
End		239	244	237	237	245	195	162
Start		43	141	45	46	45	46	52
Chain ID		V				¥	V V	
PDB UD		Iĝi	Inkr	Inkr	Inkr	lvca	lvca	lzxq
SEQ	ÖŽ	475	475	475	475	475	475	475

PDB annotation	IMMUNE SYSTEM PS8 NATURAL KILLER CELL RECEPTOR; KIR, NATURAL KILLER RECEPTOR, INHIBITORY RECEPTOR, IMMUNOGLOBULN	IMMUNE SYSTEM PS8 NATURAL KILLER CELL RECEPTOR; KIR, NATURAL KILLER RECEPTOR, INHIBITORY RECEPTOR, 2 IMMUNOGLOBULIN	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETAI; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION 3 FACTOR	COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX (TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2 NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION
Coumpound	MHC CLASS I NK CELL RECEPTOR PRECURSOR; CHAIN: A;	MHC CLASS I NK CELL RECEPTOR PRECURSOR; CHAIN: A;	GA BINDING PROTEIN ALPHA; CHAIN: A; GA BINDING PROTEIN BETA I; CHAIN: B; DNA; CHAIN: D, E;	GA BINDING PROTEIN ALPHA; CHAIN: A; GA BINDING PROTEIN BETA 1; CHAIN: B; DNA; CHAIN: D, E;	GA BINDING PROTEIN ALPHA; CHAIN: A; GA BINDING PROTEIN BETA I; CHAIN: B; DNA; CHAIN: D, E;
SeqFold Score					61.05
PMF Score	1.00	1.00	0.99	1.00	
Verify Score	0.20	0.65	0.38	0.50	
PSI BLAST Score	8.1e-18	5.4e-40	5.4e-23	1.16-21	5.4e-23
End	250	236	121	157	152
Start AA	139	45	13	13	_
Chain ID	Ą	∢	æ	æ	æ
PDB ID	2dli	2dli	lawc	lawc	lawc
SEQ ID NO:	475	475	478	478	478

	_	, ——																		
PDB annotation	3 FACTOR	TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF	TUMOR SUPPRESSOR TUMOR SUPPRESSOR, CDK4/6 INHIBITOR, ANKYRIN MOTIF	COMPLEX (KINASE/ANTI- ONCOGENE) CDK6; P16NK4A, MTS1; CYCLIN DEPENDENT KINASE	CYCLIN DEPENDENT KINASE INHIBITORY 2 PROTEIN, CDK, INK4,	CELL CYCLE, MULTIPLE TUMOR SUPPRESSOR, 3 MTS1, COMPLEX	(KINASE/ANTI-ONCOGENE) HEADER COMPLEY (KINASE/ANTI-	ONCOGENE) CDK6; P16INK4A, MTS1; CYCLIN DEPENDENT KINASE.	CYCLIN DEPENDENT KINASE	INHIBITORY 2 PROTEIN, CDK, INK4,	SUPPRESSOR, 3 MTS1, COMPLEX	(KINASE/ANTI-ONCOGENE) HEADER	COMPLEX (INHIBITOR	PROTEINKINASE) INHIBITOR PROTEIN CYCI IN. DEPENDENT	KINASE, CELL CYCLE 2 CONTROL.	ALPHA/BETA, COMPLEX (INHIBITOR	PROTEIN/KINASE)	PROTEIN/KINASE) INHIBITOR	PROTEIN, CYCLIN-DEPENDENT	KINASE, CELL CYCLE 2 CONTROL, ALPHA/BETA, COMPLEX (INHIBITOR
Coumpound		P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	P19INK4D CDK4/6 INHIBITOR; CHAIN: NULL;	CYCLIN-DEPENDENT KINASE 6; CHAIN: A; MULTIPLE TUMOR	SUPPRESSOR; CHAIN: B;		CYCLIN-DEPENDENT	KINASE 6; CHAIN: A; MULTIPLE TUMOR	SUPPRESSOR; CHAIN: B;				CYCLIN-DEPENDENT	KINASE 6; CHAIN: A; P19INK4D: CHAIN: B:			CVCI IN DEPENDENT	KINASE 6; CHAIN: A;	P19INK4D; CHAIN: B;	
SeqFold Score			56.30	52.64													15 95			
PMF Score		1.00					1.00			-			0.99							:
Verify Score		0.22					90.0		-				0.01							
PSI BLAST Score		8.1e-23	8.1e-23	2.2e-21			2.2e-21						1.4e-24			·	1 4e-24			
End AA		157	155	125			123						157		-		158		_	
Start AA		91	-	I			20						9							
Chain ID				В			В					,	X	-	_		m			
PDB ID		1bd8	1bd8	1bi7			1bi7						XIQI				15 XX		_	
SEQ NO:		478	478	478			478					1	4/8				478			

	_	,										_				_										_		
PDB annotation	PROTEIN/KINASE)	HORMONE/GROWTH FACTOR P18-	P18INK4C, TUMOR, SUPPRESSOR,	CYCLIN- 2 DEPENDENT KINASE,	HORMONE/GROWTH FACTOR	INK4C; CELL CYCLE INHIBITOR.	P18INK4C, TUMOR, SUPPRESSOR,	CYCLIN- 2 DEPENDENT KINASE,	HORMONE/GROWTH FACTOR	SIGNALING PROTEIN HELIX-TURN- HELIX ANKYRIN REPEAT		CELL CYCLE INHIBITOR P18-	INK4C(INK6); CELL CYCLE	INHIBITOR, P18-INK4C(INK6),	ANKYRIN REPEAT, 2 CDK 4/6	INHIBITOR	CELL CYCLE INHIBITOR P18- INK4C(INK6): CFL J. CYCL F	INHIBITOR, P18-INK4C(INK6).	ANKYRIN REPEAT, 2 CDK 4/6	ANK-REPEAT MYOTROPHIN	ACETYLATION, NMR, ANK-REPEAT	COMPLEX (ANTI-	ONCOGENE/ANK YRIN REPEATS)	P53BP2; ANKYRIN REPEATS, SH3, P53,	TUMOR SUPPRESSOR, MULTIGENE 2	FAMILY, NUCLEAR PROTEIN,	PHOSPHORYLATION, DISEASE	MUTATION, 3 POLYMORPHISM, COMPLEX (ANTI-
Coumpound		CYCLIN-DEPENDENT KINASE 6 INHIBITOR:	CHAIN: A;		CVCI IN. DEPENDENT	KINASE 6 INHIBITOR;	CHAIN: A;			CYCLIN-DEPENDENT KINASE 4 INHIBITOR B:	CHAIN: A;	CYCLIN-DEPENDENT	KINASE 6 INHIBITOR;	CHAIN: A, B;			CYCLIN-DEPENDENT KINASE 6 INHIBITOR:	CHAIN: A, B;		MYOTROPHIN: CHAIN:	NULL	P53; CHAIN: A; 53BP2;	CHAIN: B;					•
SeqFold Score					29 54		•										62.69											
PMF Score		1.00		_						0.77		1.00								0.82		0.64						
Verify Score		0.11							1	0.31		0.24							-	0.17		-0.01						i
PSI BLAST Score		2.4e-22			2.4e-22	 			;	5.4e-23		1.6e-22				,	1.6e-22			2.7e-21		8.1e-21						
End AA		123			161					124		123				1	159			119		128						
Start AA		13			_				5	13		13				1	-			13		13						
Chain 1D		∢			A					∢		⋖					⋖					В						
PDB ID		1bu9			1bu9				9	1098		lihb					qui			lmyo		lycs						
SEQ ID NO:		478			478				120	8/4		478				Ę	8/4			478		478						

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
										ONCOGENE/ANKYRIN REPEATS)
480	2dld	∢	18	111	5.4e-09	0.58	0.80		D-LACTATE DEHYDROGENASE; 2DLD 5 CHAIN: A, B; 2DLD 6	OXIDOREDUCTASE (CHOH(D)- NAD+(A)) R-LACTATE DEHYDROGENASE; 2DLD 7
	•									
482	1be4	V	251	308	0.00054	0.02	0.89		NUCLEOSIDE DIPHOSPHATE TRANSFERASE; CHAIN: A, B.C.	PHOSPHOTRANSFERASE PHOSPHOTRANSFERASE
482	lehw	4	251	308	0.00054	0.20	0.87		NUCLEOSIDE DIPHOSPHATE KINASE; CHAIN: A, B;	TRANSFERASE NDPK H4; NUCLEOSIDE DIPHOSPHATE KINASE, NM23, MITOCHONDRIAL, KILLER- 2 OF-PRUNE
Ş	1711	٤	0.50	900	7,0000	1	6		The state of the state of the	מוטון ז- זס
704	Inn	4	007	308	0.00034	0.31	8/:0		PHOSPHOTRANSFERASE	
									NOCLEOSIDE DIBLOSBITATE VINIASE	
									G C 2 7 4 6) COMPI EYED	
									WITH INHK 3 5'-CYCLIC	
									ADENOSINE	
									MONOPHOSPHATE INHK	
									4	
482	Inpk		250	308	0.00054	0.13	1.00		PHOSPHOTRANSFERASE(P O4 AS ACCEPTOR)	
									NUCLEOSIDE	
								_	DIPHOSPHATE KINASE	
									(E.C.2.7.4.6) INPK 3	
482	Insd	A	251	308	0.00081	-0.06	0.77		PHOSPHOTRANSFERASE	
			-						NUCLEOSIDE	
_					•			-	DIPHOSPHATE KINASE	
									(E.C.2.7.4.6) INSQ 3	
482	Inue	∀	251	310	0.00027	-0.31	0.09		NUCLEOSIDE DIPHOSPHATE KINASE:	PHOSPHOTRANSFERASE NICLEOSIDE TRIPHOSPHATE
									INUE 4 CHAIN: A, B, C, D,	NUCLEOSIDE DIPHOSPHATE INUE 10

	_	_			 				
PDB annotation			LIPID TRANSPORT APO A-I; LIPOPROTEIN, LIPID TRANSPORT, CHOLESTEROL METABOLISM, 2 ATHEROSCLEROSIS, HDL, LCAT- ACTIVATION	STRUCTURAL PROTEIN TWO REPEATS OF SPECTRIN, ALPHA HELICAL LINKER REGION, 2 2 TANDEM 3-HELIX COILED-COILS, STRUCTURAL PROTEIN	RNA-BINDING PROTEIN/RNA TRA PRE-MRNA; SPLICING REGULATION, RNP DOMAIN, RNA COMPLEX		GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN I, PABP I; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM,
Coumpound	E, F; 1NUE 5		APOLIPOPROTEIN A-1; CHAIN: A, B, C, D;	ALPHA SPECTRIN; CHAIN: A, B, C;	SXL-LETHAL PROTEIN; CHAIN: A, B; RNA (5: R(P*GP*UP*UP*GP*UP*UP *In*In*In*In*In*In*In*In*In*In*In*In*In*	CHAIN: P, Q;	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP*AP* AP*AP*AP*AP*AP* CHAIN: M, N, O, P, Q, R, S, T.	POLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B, C, D, E, F, G, H; RNA (5- R(*AP*AP*AP*AP*AP* AP*AP*AP*AP*AP* CHAIN: M, N, O, P, Q, R, S,	PÓLYDENYLATE BINDING PROTEIN 1; CHAIN: A, B,
SeqFold Score			50.39						
PMF Score				0.06	0.13		0.07	0.31	0.03
Verify Score				-0.43	0.07		0.44	0.25	0.28
PSI BLAST Score			0.00081	2.7e-05	0.00014		0.0027	0.0027	0.0027
End AA			174	150	134		134	134	134
Start AA				11	48		48	48	48
Chain ID			∢	4	4		4	Ф	ţ L
PDB ID			lavi	lcun	1b7f		lcvj	lcvj	lcvj
SEQ ID NO:			483	483	484		484	484	484

S E S	PDB ID	Chain ID	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP*AP* AP*AP*AP*AP*A)-3'); CHAIN: M, N, O, P, Q, R, S, T:	PROTEIN-RNA COMPLEX, GENE REGULATION/RNA
484	lcvj	н	48	134	0.0027	0.68	0.39		POLYDENYLATE BINDING PROTEIN I; CHAIN: A, B, C, D, E, F, G, H; RNA (5'- R(*AP*AP*AP*AP*AP* AP*AP*AP*A,3'); CHAIN: M, N, O, P, Q, R, S, T;	GENE REGULATION/RNA POLY(A) BINDING PROTEIN 1, PABP 1; RRM, PROTEIN-RNA COMPLEX, GENE REGULATION/RNA
484	1482	Ą	48	134	0.0011	0.33	0.05		HU ANTIGEN C; CHAIN: A;	RNA BINDING PROTEIN RNA- BINDING DOMAIN
484	2u2f	A	48	134	0.00027	0.43	0.27		SPLICING FACTOR UZAF 65 KD SUBUNIT; CHAIN: A;	RNA-BINDING PROTEIN SPLICING, U2 SNRNP, RBD, RNA-BINDING PROTEIN
1										
485	1d0s	∢	75	217	1.6e-09	0.13	-0.20		NICOTINATE MONONUCLEOTIDE:5,6- CHAIN: A;	TRANSFERASE DINUCLEOTIDE- BINDING MOTIF, PHOSPHORIBOSYL TRANSFERASE
485	1req	4	16	193	2.7e-13	0.33	-0.19		METHYLMALONYL-COA MUTASE; CHAIN: A, B, C, D;	ISOMERASE ISOMERASE, MUTASE, INTRAMOLECULAR TRANSFERASE
485	1req	A	17	227	8.1e-10	0.04	-0.20		METHYLMALONYL-COA MUTASE; CHAIN: A, B, C, D;	ISOMERASE ISOMERASE, MUTASE, INTRAMOLECULAR TRANSFERASE
	1aab		2	84	2.2e-30	1.08	1.00		HIGH MOBILITY GROUP PROTEIN; 1AAB 5 CHAIN: NULL; 1AAB 6	DNA-BINDING HMGA DNA-BINDING HMG-BOX DOMAIN A OF RAT HMG1; 1AAB 8 HMG-BOX 1AAB 20
489	1aab		2	84	2.2e-30			131.01	HIGH MOBILITY GROUP PROTEIN; 1AAB 5 CHAIN: NULL; 1AAB 6	DNA-BINDING HMGA DNA-BINDING HMG-BOX DOMAIN A OF RAT HMG1; 1AAB 8 HMG-BOX 1AAB 20

	1						
PDB annotation	DNA BINDING PROTEIN HMG BOX, DNA BENDING, DNA RECOGNITION, CHROMATIN, NMR, DNA 2 BINDING PROTEIN	DNA BINDING PROTEIN HMG BOX, DNA BENDING, DNA RECOGNITION, CHROMATIN, NMR, DNA 2 BINDING PROTEIN	GENE REGULATION/DNA HMG-1, AMPHOTERIN, HEPARIN-BINDING PROTEIN P30; HIGH-MOBILITY GROUP DOMAIN, BENT DNA, PROTEIN-DRUG-DNA 2 COMPLEX, GENE REGULATION/DNA	GENE REGULATION/DNA HMG-1, AMPHOTERIN, HEPARIN-BINDING PROTEIN P30; HIGH-MOBILITY GROUP DOMAIN, BENT DNA, PROTEIN-DRUG-DNA 2 COMPLEX, GENE REGULATION/DNA	ENDOCYTOSIS/EXOCYTOSIS SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX BUNDLE	ISOMERASE ISOMERASE, MUTASE, INTRAMOLECULAR TRANSFERASE	COMPLEX (TRANSDUCER/TRANSDUCTION) GT BETA-GAMMA; MEKA, PP33; PHOSDUCIN, TRANSDUCIN, BETA- GAMMA, SIGNAL TRANSDUCTION, 2 REGULATION, PHOSPHORYLATION, G PROTEINS, THIOREDOXIN, 3
Coumpound	NON HISTONE PROTEIN 6 A; CHAIN: A;	NON HISTONE PROTEIN 6 A; CHAIN: A;	HIGH MOBILITY GROUP I PROTEIN; CHAIN: A; DNA (5'-D(*CP*CP*(IDO) CHAIN: B; DNA (5'- CHAIN: C;	HIGH MOBILITY GROUP 1 PROTEIN; CHAIN: A; DNA (5'-D(*CP*CP*(IDO) CHAIN: B; DNA (5'- CHAIN: C;	SYNTAXIN-1A; CHAIN: A, B, C;	METHYLMALONYL-COA MUTASE; CHAIN: A, B, C, D;	TRANSDUCIN; CHAIN: B, G; PHOSDUCIN; CHAIN: P;
SeqFold Score	77.91			106.80			
PMF Score		1.00	1.00		-0.19	-0.20	-0.18
Verify Score		0.59	0.80		0.35	0.28	0.13
PSI BLAST Score	1.1e-29	1.1e-29	1.16-24	1.1e-24	2.7e-08	2.2e-08	2.7e-10
End AA	167	165	78	78	211	211	218
Start	75	.63	∞	∞	145	145	145
Chain ID	A	A	٨	Ą	A	¥	۵.
PDB ID	1cg7	1cg7	1ckt	Ickt	lez3	Ireq	2trc
SEQ ID NO:	489	489	489	489	489	489	489

		_							,																	
PDB annotation	VISION, MEKA, COMPLEX (TRANSDUCENTRANSDUCTION)		COMPLEX (TRANSCRIPTION REGULATION/DNA) GABPALPHA; GABPBETA1; COMPLEX	(TRANSCRIPTION REGULATION/DNA), DNA-BINDING, 2	NUCLEAR PROTEIN, ETS DOMAIN, ANKYRIN REPEATS, TRANSCRIPTION	3 FACTOR	SIGNAL TRANSDUCTION SIGNAL	HOMOLOGY (PH) DOMAIN	COMPLEX (DNA-BINDING	PROTEIN/DNA) SERUM RESPONSE	FACTOR ACCESSORY PROTEIN 1A;	EIS DOMAIN, DNA-BINDING	DOMAIN, WINGED HELIX-TURN-	HELIX, 2 CKYSTAL STRUCTURE,	DINA-BINDING SPECIFICITY,	COMPLEX 3 (DNA-BINDING	PROTEIN/DNA) SHEET HEADER	TRANSFERASE BRUTON'S	AGAMMAGLOBULINEMIA TYROSINE	KINASE, BTK; TRANSFERASE, PH	DOMAIN, BTK MOTIF, ZINC BINDING,	X-LINKED 2	AGAMMAGLOBULINEMIA,	TYROSINE-PROTEIN KINASE	GENE REGULATION SON OF	SEVENLESS PROTEIN; GUANINE NUCLEOTIDE EXCHANGE FACTOR,
Coumpound			GA BINDING PROTEIN ALPHA; CHAIN: A; GA BINDING PROTEIN BETA	I; CHAIN: B; DNA; CHAIN: D, E;			SOS1; CHAIN: NULL;		E74 PROMOTOR DNA;	CHAIN: A, B; SAP-1;	CHAIN: C;							BRUTON'S TYROSINE	KINASE; CHAIN: A, B;						HUMAN SOS 1; CHAIN: A;	
SeqFold Score														-		-										
PMF Score			0.42				0.19		0.28									0.89							0.40	
Verify Score			-0.22				-0.04		-0.30									-0.33							-0.07	-
PSI BLAST Score			0.0054	**		_	0.00054		0.0054									0.0011						1	0.00027	
End AA			134				Ξ		134									107							109	
Start AA			92				47		06									34							55	_
Chain 1D			∢						၁									A							Α	
PDB ID			lawc				lawe		1bc8									1btk					•		ldbh	
SEQ NO:			490				490		490									490							490	

SEQ ID NO:	PDB ID	Chain ID	Start	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
										GENE REGULATION
490	1dux	O	16	134	0.0027	-0.08	0.22		DNA (5'- D(*TP*GP*AP*CP*CP*GP* GP*AP*AP*GP*TP*GP*T)	TRANSCRIPTION/DNA TRANSFORMING PROTEIN ELK-1, ELK1, MEMBER OF ETS ETS-DOMAIN,
									CHAIN: A, D; DNA (5- D(*AP*CP*AP*CP*TP*TP*	DNA-BINDING DOMAIN, WINGED HELIX-TURN-HELIX, 2 CRYSTAL
									CP*CP*GP*IP*CP*A) CHAIN: B, E; ETS-DOMAIN PROTEIN ELK-1; CHAIN: C, F;	STRUCTURE, DNA-BINDING SPECIFICITY
490	letc		8	134	0.0027	-0.47	0.09		MURINE ETS-1 TRANSCRIPTION FACTOR; IETC 4 CHAIN: NULL; IETC 5	TRANSCRIPTION REGULATION
490	1fao	⋖	78	106	2.2e-13	0.73	1.00		DUAL ADAPTOR OF PHOSPHOTYROSINE AND	SIGNALING PROTEIN DAPP1, PHISH, BAM32: PLECKSTRIN. 3-
									3- CHAIN: A;	PHOSPHOINOSITIDES, INOSITOL TETRAKISPHOSPHATE 2 SIGNAL
										TRANSDUCTION PROTEIN, ADAPTOR PROTEIN
490	1fb8	A	28	901	5.4e-13	0.92	1.00		DUAL ADAPTOR OF PHOSPHOTYROSINE AND	SIGNALING PROTEIN DAPP1, PHISH, BAM32-PI ECKSTRIN 3.
									3- CHAIN: A;	PHOSPHOINOSITIDES, INOSITOL
										TRANSDUCTION PROTEIN, ADAPTOR PROTEIN
490	lfgy	V	28	901	5.4e-12	0.91	1.00		GRP1; CHAIN: A;	SIGNALING PROTEIN ARFI GUANINE NUCLEOTIDE EXCHANGE FACTOR AND PH DOMAIN
490	ig:	Ą	08	135	0.0027	-0.07	0.09		FLI-1; IFLI 5 CHAIN: A; IFLI 6 DNA IFLI 10 CHAIN: B, C; IFLI 12	COMPLEX (TRANSCRIPTION FACTOR/DNA)
490	1pls		28	106	5.4e-11	0.29	00:1		PHOSPHORYLATION PLECKSTRIN (N.	

Chain ID	iii C	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation	
								HOMOLOGY DOMAIN) MUTANT IPLS 3 WITH LEU GLU (HIS)6 ADDED TO THE C TERMINUS IPLS 4 (INS(G105-LEHHFHHHF)) (NMR, 25 STRUCTURES) IPLS 5		
		28	801	2.7e-10	0.12	86.0		SOS 1; CHAIN: NULL;	SIGNAL TRANSDUCTION SON OF SEVENLESS; PLECKSTRIN, SON OF SEVENLESS, SIGNAL TRANSDUCTION	
∢ .		10	212	3.4e-41	0.20	1.00		ULPI PROTEASE; CHAIN: A; UBITQUTIN-LIKE PROTEIN SMT3; CHAIN: B;	HYDROLASE SUMO HYDROLASE, UBIQUITIN-LIKE PROTEASE 1, SMT3 HYDROLASE 2 DESUMOYLATING ENZYME, CYSTEINE PROTEASE, SUMO PROCESSING 3 ENZYME, SMT3 PROCESSING ENZYME, NABH4, THIOHEMIACETAL, 4 COVALENT PROTEASE ADDUCT	
В		179	323	8.1e-09	0.02	-0.14		SYNTAXIN BINDING PROTEIN 1; CHAIN: A; SYNTAXIN 1A; CHAIN: B;	ENDOCYTOSIS/EXOCYTOSIS NSEC1; PROTEIN-PROTEIN COMPLEX, MULTI- SUBUNIT	_
A		31	226	0.00017			61.44	APOLIPOPROTEIN A-1; CHAIN: A, B, C, D;	LIPID TRANSPORT APO A-I; LIPOPROTEIN, LIPID TRANSPORT, CHOLESTEROL METABOLISM, 2 ATHEROSCLEROSIS, HDL, LCAT- ACTIVATION	
A		40	245	8.1e-07			54.20	ALPHA SPECTRIN; CHAIN: A, B, C;	STRUCTURAL PROTEIN TWO REPEATS OF SPECTRIN, ALPHA HELICAL LINKER REGION, 2 2 TANDEM 3-HELIX COILED-COILS,	

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
										STRUCTURAL PROTEIN
501	Iquu	Ą	20	243	2.7e-07			55.43	HUMAN SKELETAL MUSCLE ALPHA-ACTININ	CONTRACTILE PROTEIN TRIPLE- HELIX COILED COIL, CONTRACTILE
									z; CHAIN: A;	PROTEIN
202	1ad0	∢	20	225	1.7e-82			101.11	FAB FRAGMENT,	IMMUNOGLOBULIN RAB ER AGMENT
									A, B, C, D;	INTERIOR OFFICE INTERIOR INTER
205	lafv	r	21	224	3.4e-78			101.19	HUMAN	COMPLEX (VIRAL
									IMMUNODEFICIENCY	CAPSID/IMMUNOGLOBULIN) HIV-1
									CHAIN: A, B; ANTIBODY	LIGHT CHAIN, FAB HEAVY CHAIN
									FAB25.3 FRAGMENT;	COMPLEX (VIRAL
									CHAIN: H, K, L, M;	CAPSID/IMMUNOGLOBULIN), HIV,
\$03	1001	,	ç	700	1 20 70	010	000		THE EAD. CHIABL. I. II.	CAPSID PROTEIN, 2 F24
300	1471	<u>-</u>	3	077	1.25-79	61.0	0.30		IF / FAB, CRAIN: L, II,	IMMINOGLOBOLIN ANTIBODY FAR
										ENZYME INHIBITOR, PCR, 2 HOT
										START
502	1b2w	1	20	229	6.8e-87	.0.15	0.84		ANTIBODY (LIGHT	IMMUNE SYSTEM
									CHAIN); CHAIN: L;	IMMUNOGLOBULIN;
									ANTIBODY (HEAVY	IMMUNOGLOBULIN ANTIBODY
									CHAIN); CHAIN: H;	ENGINEERING, HUMANIZED AND
										CHIMERIC ANTIBODY, FAB, 2 X-RAY
					-					SIRUCIUKE, IHKEE-DIMENSIUNAL
										SIKYCIUKE, GAMMA- 3
\$02	1h2w	1	2	27.5	6 80-87			100 05	ANTIBODY (1 IGET	INTERFERON, IMMUNE SYSTEM
!		1	i .	ì	3			6000	CHARD: CHARD: 1.	MANAGE STEERS
	_								ANTIBODY (HEAVY	IMMENOGEOBOLIN,
									CHAIN: CHAIN: H:	ENGINEERING, HUMANIZED AND
										CHIMERIC ANTIBODY, FAB, 2 X-RAY
										STRUCTURE, THREE-DIMENSIONAL
										STRYCTURE, GAMMA-3

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PDB annotation	INTERFERON, IMMUNE SYSTEM	IMMUNOGLOBULIN IMMUNOGLOBULIN, KAPPA LIGHT- CHAIN DIMER HEADER	IMMUNOGLOBULIN IMMUNOGLOBULIN, KAPPA LIGHT- CHAIN DIMER HEADER		COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA A2 HEAVY CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR)	ANTIBODY ANTIBODY, FAB, CAMPATH-1G, CD52	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	ANTIBODY THERAPEUTIC, ANTIBODY, CD52
Coumpound		IMMUNOGLOBULIN; CHAIN: A, B;	IMMUNOGLOBULIN; CHAIN: A, B;	IMMUNOGLOBULIN FAB' FRAGMENT OF MONOCLONAL ANTIBODY B72.3 IBBJ 3 (MURINE/HUMAN CHIMERA) IBBJ 4	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E:	CAMPATH-1G ANTIBODY; CHAIN: A, B, C, D, E, F, G, H;	FAB FRAGMENT; CHAIN: L, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR; CHAIN: V, W;	FAB FRAGMENT; CHAIN: L, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR; CHAIN: V, W;	CAMPATH-1H:LIGHT CHAIN; CHAIN: L;
SeqFold Score			109.72	102.84	233.42	105.12		109.40	
PMF Score		96.0					0.90		0.99
Verify Score		0.11					0.03		0.27
PSI BLAST Score		1.4e-86	1.4e-86	1.4e-78	8.1e-74	5.1e-79	1.4e-88	1.4e-88	6.8e-85
End		226	234	233	219	235	228	224	226
Start AA		20	21	21	21	21	20	21	20
Chain ID		A	Y	ว	Ω	А	T	ſ	L
PDB ID		156d	156d	1 b bj	1642	1bfo	16j1	16j1	lcel
SEQ ID NO:		502	502	502	502	502	502	502	502

PDB annotation			ANTIBODY, CD52 ANTIBODY, CD52	A, TRANSITION STATE ANALOG, IMMUNE SYSTEM	CRYSTAL STRUCTURE 2.7A CRYSTAL STRUCTURE 2.7A RESOLUTION BINDING 2 OUTSIDE THE ANTIGEN COMBINING SITE SUPERANTIGEN FAB VH3 3 SPECIFICITY		N IMMUNE SYSTEM IMMUNOGLOBULIN, ANTIBODY, FAB, HEPATITIS B, PRES2 G1 B,		C- IMMUNE SYSTEM VON WILLEBRAND
Coumpound		CAMPATH-IH:HEAVY CHAIN; CHAIN: H; PEPTIDE ANTIGEN; CHAIN: P;	CAMPATH-IH:LIGHT CHAIN; CHAIN: L; CAMPATH-IH:HEAVY CHAIN; CHAIN: H; PEPTIDE ANTIGEN; CHAIN: P;	7C8 FAB FRAGMENT; SHORT CHAIN; CHAIN: A, C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D	IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B, D, F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H;	IMMUNOGLOBULIN 3D6 FAB 1DFB 3	F124 IMMUNOGLOBULIN (KAPPA LIGHT CHAIN); CHAIN: A, C; F124 IMMUNOGLOBULIN (IGG1 HEAVY CHAIN); CHAIN: B, D;	IMMUNOGLOBULIN IMMUNOGLOBULIN G1 (KAPPA LIGHT CHAIN) FAB' FRAGMENT 1FIG 3	IMMUNOGLOBULIN NMC-
SeqFold	31036		108.57	105.20		107.04		103.10	
PMF	2006			·	0.95		1.00		0.77
Verify	31000				0.06		0.05		0.18
PSI BI AST	Score		6.8e-85	5.1e-75	1.2e-89	le-84	1.7e-79	5.1e-81	1.7e-85
End	nn I		233	225	229	225	226	225	229
Start	V V		21	21	20	21	20	20	20
Chain			1	∢	∢	7	٧	1	ı
EDB EDB	2		IceI	lct8	Idee	1dfb	1f11	lfig	lfns
SEQ	Ş Ş		502	502	502	502	502	502	202

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PDB annotation	(A:ALPHA) BINDING, 2 COMPLEX (WILLEBRAND/IMMUNOGLOBULIN), BLOOD COAGULATION TYPE 3 2B VON WILLEBRAND DISEASE			COMPLEX (HIV ENVELOPE PROTEIN/CD4FAB) COMPLEX (HIV ENVELOPE PROTEIN/CD4/FAB), HIV-1 EXTERIOR 2 ENVELOPE GP120, T- CELL SURFACE GLYCOPROTEIN CD4, 3 ANTIGEN-BINDING FRAGMENT OF HUMAN IMMUNOGLOBULIN 17B, 4 GLYCOSYLATED PROTEIN	COMPLEX (HIV ENVELOPE PROTEIN/CD4/FAB) COMPLEX (HIV ENVELOPE PROTEIN/CD4/FAB), HIV-1 EXTERIOR 2 ENVELOPE GP120, T- CELL SURFACE GLYCOPROTEIN CD4, 3 ANTIGEN-BINDING FRAGMENT OF HUMAN IMMUNOGLOBULIN 17B, 4 GLYCOSYLATED PROTEIN		
Coumpound	IMMUNOGLOBULIN NMC- 4 IGGI; CHAIN: H; VON WILLEBRAND FACTOR; CHAIN: A:	IMMUNOGLOBULIN FAB FRAGMENT OF HUMANIZED ANTIBODY 4DS, VERSION 4 IFVD 3	IMMUNOGLOBULIN FAB FRAGMENT OF HUMANIZED ANTIBODY 4D5, VERSION 4 1FVD 3	ENVELOPE PROTEIN GP120; CHAIN: G; CD4; CHAIN: C; ANTIBODY 17B; CHAIN: L, H;	ENVELOPE PROTEIN GP120; CHAIN: G; CD4; CHAIN: C; ANTIBODY 17B; CHAIN: L, H;	IMMUNOGLOBULIN IGG2A FAB FRAGMENT (FAB 17/9) IHIL 3	IMMUNOGLOBULIN IGG2A FAB FRAGMENT (FAB 17/9) COMPLEX
SeqFold Score			106.39	108.90			
PMF Score		86.0			1.00	0.90	0.81
Verify Score		0.08			0.17	0.14	0.00
PSI BLAST	Score	1.4e-86	1.4e-86	1,5e-81	1.5e-81	5.1e-81	5.1e-81
End AA		229	225	233	226	226	226
Start AA		20	21	21	23	20	20
Chain ID		A	¥	7		A	J.
PDB CD		1fvd	1fvd	1gc1	lgcl	lhil	1jih
SEQ EQ		502	502	. 502	502	202	502

SEQ ID NO:	PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									WITH PEPTIDE OF 11FH 3 INFLUENZA HEMAGGLUTININ HA1 (STRAIN X47) (RESIDUES	
502	ligt	A	70	229	3.4e-85	0.14	0.90		IGG2A INTACT ANTIBODY - MAB231; CHAIN: A, B, C, D	IMMUNOGLOBULIN INTACT IMMUNOGLOBULIN V REGION C REGION, IMMUNOGLOBULIN
502	ligt	٧	21	225	3.4e-85			102.74	IGG2A INTACT ANTIBODY - MAB231; CHAIN: A, B, C, D	IMMUNÓGLOBULIN INTACT IMMUNOGLOBULIN V REGION C REGION, IMMUNOGLOBULIN
502	1kb5	7	21	225	1.7e-83			108.53	KB5-C20 T-CELL ANTIGEN RECEPTOR; CHAIN: A, B; ANTIBODY DESIRE-1; CHAIN: I, H;	COMPLEX (IMMUNOGLOBULIN/RECEPTOR) TCR VAPLHA VBETA DOMAIN; T-CELL RECEPTOR, STRAND SWITCH, FAB, ANTICLONOTYPIC, 2 (IMMUNOGLOBULIN/RECEPTOR)
502	lnsn	ы	20	228	3.46-81	0.10	98.0		IGG FAB (IGGI, KAPPA); INSN 4 CHAIN: L, H; INSN 5 STAPHYLOCOCCAL NUCLEASE; INSN 9 CHAIN: S; INSN 10	COMPLEX (IMMUNOGLOBULIN/HYDROLASE) NI0 FAB IMMUNOGLOBULIN; INSN 7 STAPHYLOCOCCAL NUCLEASE RIBONUCLEATE, INSN 11 IMMUNOGLOBULIN, STAPHYLOCOCCAL NUCLEASE INSN 25
202	lqrn	Ω	21	222	2.4e-68			251.37	MHC CLASS I HLA-A; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE P6A; CHAIN: C; HMAN T-CELL RECEPTOR; CHAIN: D; HLA-A 0201: CHAIN: E:	IMMUNE SYSTEM HUMAN TCR/PEPTIDE/MHC COMPLEX, HLA- A2, HTLV-1, TAX, TCR, T2 CELL RECEPTOR, IMMUNE SYSTEM
505	1sbs	J	70	229	5.1c-85	0.20	96.0		MONOCLONAL ANTIBODY 3A2; CHAIN: H,	MONOCLONAL ANTIBODY MONOCLONAL ANTIBODY, FAB-

SEQ NO:	PDB CI	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									L,	FRAGMENT, REPRODUCTION
502	1tcr	A	21	227	1.4e-74			181.32	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL
502	lvge	1	21	225	1.7e-85			106.15	TRI.9 FAB; CHAIN: L, H;	IMMUNOGLOBULIN TRI.9, ANTI- THYROID PEROXIDASE, AUTOANTIBODY, 2 IMMUNOGLOBULIN
502	lvge	7	22	229	1.7e-85	0.05	0.87		TRI.9 FAB; CHAIN: L, H;	IMMUNOGLOBULIN TRI.9, ANTI- THYROID PEROXIDASE, AUTOANTIBODY, 2 IMMUNOGLOBULIN
502	25c8	Г	20	226	5.1e-81	0.04	0.72		IGG 5C8; CHAIN: L, H;	CATALYTIC ANTIBODY CATALYTIC ANTIBODY, FAB, RING CLOSURE REACTION
502	2fgw	J	20	229	3.4e-89	0.09	56'0		IMMUNOGLOBULIN FAB FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 1452' (HUH52- OZ FAB) 2FGW 4	·
502	2fgw	ı	21	225	3.46-89			111.79	IMMUNOGLOBULIN FAB FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4	
502	3fct	A	22	228	5.1e-8]	0.20	0.83		METAL CHELATASE CATALYTIC ANTIBODY; CHAIN: A, C; METAL CHELATASE CATALYTIC ANTIBODY; CHAIN: B, D;	IMMUNE SYSTEM METAL CHELATASE, CATALYTIC ANTIBODY, FAB FRAGMENT, IMMUNE 2 SYSTEM
203	lao7	E	22	180	2.7e-49	0.29	1.00		HLA-A 0201; CHAIN: A;	COMPLEX (MHC/VIRAL

PDB annotation	JILN; PEPTIDE/RECEPTOR) HLA-A2 HEAVY DE; CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR	;; COMPLEX (MHC/VIRAL ILIN; PEPTIDE/RECEPTOR) HLA-A2 HEAVY DE; CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR	GOMPLEX (MHC/VIRAL ILIN; PEPTIDE/RECEPTOR) HLA A2 HEAVY DE; CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) NIN:	; COMPLEX (MHC/VIRAL ILIN; PEPTIDE/RECEPTOR) HLA A2 HEAVY DE; CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) NIN:	N RECEPTOR T CELL RECEPTOR IBEC 14
Coumpound	BETA-2 MICROGLOBULN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN:	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	14.3.D T CELL ANTIGEN RECEPTOR; IBEC 5 CHAIN: NITT: IBEC 6
SeqFold Score		132.96		147.96	135.25
PMF Score			1.00		
Verify Score			0.48		
PSI BLAST Score		2.7e-49	5.4e-54	5.4e-54	5.4e-51
End		180	180	180	081
Start AA		22	22	22	22
Chain ID		ம	កា	ம	
PDB ID		1ao7	16d2	1642	1 bec
SEQ ID NO:		203	503	503	503

PDB annotation		RECEPTOR T CELL RECEPTOR IBEC	IMMUNE SYSTEM HLA-DRI, DRA; HLA-DRI, DRBI 0101; TCR HA1.7 ALPHA CHAIN; TCR HA1.7 BETA CHAIN; PROTEIN-PROTEIN COMPLEX, IMMUNOGLOBULIN FOLD			SIGNAL TRANSDUCTION PROTEIN	CYTOSKELETON	SH3 PROTOTYPE WWPROTOTYPE, PROTEIN DESIGN	ISOMERASE PINI; PEPTIDYL- PROLINE ISOMERASE, WW DOMAIN, PHOSPHOSERINE BINDING	SIGNALING PROTEIN DAPPI, PHISH, BAM32; PLECKSTRIN, 3-PHOSPHOINOSITIDES, INOSITOL TETRAKISPHOSPHATE 2 SIGNAL TRANSDUCTION PROTEIN, ADAPTOR	SIGNALING PROTEIN DAPPI, PHISH, BAM32; PLECKSTRIN, 3-
Coumpound		14.3.D T CELL ANTIGEN RECEPTOR; IBEC 5 CHAIN: NULL; IBEC 6	HLA CLASS II HISTOCOMPATBILITY ANTIGEN, DR CHAIN: A; HLA CLASS II HISTOCOMPATBILITY ANTIGEN, DR-1 CHAIN: B; HEMAGGLUTININ HAI	PET ILDE CHANN; CHAIN: C; T-CELL RECEPTOR ALPHA CHAIN; CHAIN: D; T-CELL RECEPTOR BETA	CHAIN; CHAIN: E;	BETA-SPECTRIN; 1BTN 4 CHAIN: NULL; 1BTN 5	90	Ë	PEPTIDYL-PROLYL CIS- TRANS ISOMERASE NIMA- CHAIN: B; Y(SEP)PT(SEP)S PEPTIDE: CHAIN: C:	AND	DUAL ADAPTOR OF PHOSPHOTYROSINE AND
SeqFold	Score			<u>-,, -, -</u>							
PMF	Score	1.00	1.00			0.88	9.76	0.25	0.07	0.94	0.82
Verify	Score	0.47	0.36			0.88	0.67	0.08	0.08	0.61	09:0
PSI	BLAS I Score	5.4e-51	5.4e-57			1.1e-11	5.4e-15	0.0014	0.00054	2.7e-11	8.1e-11
End	¥¥	180	180			246	251	79	59	253	253
Start	AA	23	23			153	153	35	35	136	136
Chain	OT .		മ					А	В		A
BOY CT	ar I	1bec	1 <u>f</u>			1btn	ldro	le0m	1f8a	lfao	1fb8
SEQ	N S S	503	503			505	505	505	505	505	505

PDB annotation	PHOSPHOINOSITIDES, INOSITOL TETRAKISPHOSPHATE 2 SIGNAL TRANSDUCTION PROTEIN, ADAPTOR PROTEIN	SIGNALING PROTEIN ARFI GUANINE NUCLEOTIDE EXCHANGE FACTOR AND PH DOMAIN	RIN H ED IPLS HH) SS)	SIGNAL TRANSDUCTION SON OF SEVENLESS; PLECKSTRIN, SON OF SEVENLESS, SIGNAL TRANSDUCTION	SIGNAL TRANSDUCTION IRS-1; BETA- 4: A, SANDWHICH, SIGNAL TRANSDUCTION				AL PROTEIN HS6: 50S RIBOSOMAL		N RIBOSOMAL PROTEIN L14P, HMAL14,
Coumpound	3- CHAIN: A;	GRP1; CHAIN: A;	PHOSPHORYLATION PLECKSTRIN (N- TERMINAL PLECKSTRIN HOMOLOGY DOMAIN) MUTANT IPLS 3 WITH LEU GLU (HIS)6 ADDED TO THE C TERMINUS IPLS 4 (INS(G105-LEHTHHHH)) (NMR, 25 STRUCTURES) IPLS 5	SOS I; CHAIN: NULL;	INSULIN RECEPTOR SUBSTRATE I; CHAIN: A, B;	o in the read over	23S KKNA; CHAIN: 0; 5S RRNA; CHAIN: 9; RIBOSOMAL PROTEIN L2; CHAIN: A; RIBOSOMAL	PROTEIN L3; CHAIN: B;	CHAIN: C: RIBOSOMAL	PROTEIN LS; CHAIN: D;	RIBOSOMAL PROTEIN
SeqFold Score											-
PMF Score		0.88	0.80	0.36	0.17	13.0	0.64		7.		
Verify Score		0.74	0.18	0.41	0.21	6	-0.29				
PSI BLAST Score		8.1e-06	2.7e-10	1.9e-14	1.9e-12		1./e-19				
End		252	250	252	315	8	33				
Start		158	136	130	138	=	=				
Chain ID		Ą			A	Ç	ਸ				
PDB ID		Ifgy	1pis	1pms	lqqg	183.	Ĭ				
SEQ ID NO:		505	505	505	505	203)00				

			SO S	, 65	.19E,		JWMC	SO	AL23,	TEIN			OTEIN)MAL	L16;	L34,	L32E,	.37E,	S	MAL	-	L6,	Ι Α -												-
ion		SOMAL	PROTEIN L18P, HMAL18, HL12; 50S	RIBOSOMAL PROTEIN LISE, HL29	L19; 50S RIBOSOMAL PROTEIN L19E,	HMAL 19, HL24; 50S RIBOSOMAL	PROTEIN L21E, HL31; 50S RIBOSOMAL	PROTEIN L22P, HMAL22, HL23; 50S	RIBOSOMAL PROTEIN L23P, HMAL23,	HL25, L21; 50S RIBOSOMAL PROTEIN	.15; 50S	,24E,	HL21/HL22; 50S RIBOSOMAL PROTEIN	L29P, HMAL29, HL33; 50S RIBOSOMAL	PROTEIN L30P, HIMAL30, HL20, HL16;	50S RIBOSOMAL PROTEIN L31E, L34,	HL30; 50S RIBOSOMAL PROTEIN L32E,	HLS; 50S RIBOSOMAL PROTEIN L37E,	L35E; 50S RIBOSOMAL PROTEINS	.39E, HL39E, HL46E; 50S RIBOSOMAL	; 50S	RIBOSOMAL PROTEIN LGP, HIMAL6,	HL10 RIBOSOME ASSEMBLY, RNA-	OTEIN-											
PDB annotation		50S RIBO	HMAL 18	COLEIN	OMAL P	50S RIB	HL31; 50	HMAL22	OTEIN 1	UBOSON	HL16, HI	COTEIN 1	RIBOSO	HL33; 50	HIMAL30	L PROTE	SOMAL 1	OMAL P	SOMAL I	.46E; 50S	LA, HLA	OTEINI	E ASSEN	RNA, PR											
P.D.		HMAL 15, HL9; 50S RIBOSOMAI	IN LI8P,	OMAL PI	S RIBOS	19, HL24;	IN L21E,	IN L22P,	OMAL PI	L21; 50S I	L24P, HMAL24, HL16, HL15; 50S	RIBOSOMAL PROTEIN L24E,	IL22; 50S	IMAL29,	IN L30P,	30SOMA	SOS RIBO	S RIBOS	OS RIBO	11.39E, HI	PROTEIN L44E, LA, HLA; 50S	OMAL PF	IBOSOM	RNA, PROTEIN-RNA, PROTEIN	Z										
		HIMAL	PROTE	RIBOS	L19; 50	HMAL	PROTE	PROTE	RIBOS	HL25,]	L24P, I	RIBOS	HL21/F	L29P, F	PROTE	50S RII	HL30;	HL5; 5(L35E;	L39E, I	PROTE	RIBOS	HL10 R	RNA, P	PROTEIN										
P		rein		LEIN		IEIN		IEIN		TEIN		IEIN		EIN		TEIN		EIN		EIN		TEIN		EIN		EIN		EIN		EIN		EIN		EIN	
Coumpound		TAL PRO	AIN: F;	IAL PRO	: :::	IAL PRO	H.H.	TAL PRO	AIN: I;	TAL PRO	Z: 7;	TAL PRO	N: K;	IAL PRO	AIN: L;	IAL PROT	N: M;	IAL PRO	AIN: N;	(AL PRO)	Ö;	(AL PRO)	R. P.	(AL PRO)	ö Z	IAL PROJ	AIN: R;	IAL PROJ	R: S;	IAL PROJ	Ä.T.	IAL PROJ	AIN: U;	AL PROT	.; N: V:
0		RIBOSOMAL PROTEIN	L10E; CHAIN: F	KIBOSOMAL PROTEIN	LI3; CHAIN: G;	RIBOSOMAL PROTEIN	L14; CHAIN: H;	RIBOSOMAL PROTEIN	LISE; CHAIN: I;	RIBOSOMAL PROTEIN	L15; CHAIN: J;	RIBOSOMAL PROTEIN	L18; CHAIN: K;	RIBOSOMAL PROTEIN	L18E; CHAIN: L;	RIBOSOMAL PROTEIN	L19; CHAIN: M;	RIBOSOMAL PROTEIN	L21E; CHAIN: N;	RIBOSOMAL PROTEIN	L22; CHAIN: 0;	RIBOSOMAL PROTEIN	L23; CHAIN: P;	RIBOSOMAL PROTEIN	L24; CHAIN: Q;	RIBOSOMAL PROTEIN	L24E; CHAIN: R;	RIBOSOMAL PROTEIN	L29; CHAIN: S;	RIBOSOMAL PROTEIN	L30; CHAIN: T;	RIBOSOMAL PROTEIN	L31E; CHAIN: U;	RIBOSOMAL PROTEIN	L32E; CHAIN: V;
SeqFold	Score						-	-			_			-									-	-										-	
PMF	Score																																		
Verify	Score																																		
PSI	Score																																		
End	AA A																																		
Start	AA								_																										
Chain	LU																																		
PDB	a																																		
SEQ	g ö				_							_																							

		_			_			·	
PDB annotation			TRYPTOPHAN BIOSYNTHESIS TRYPTOPHAN INDOLE-LYASE; TRYPTOPHAN BIOSYNTHESIS, TRYPTOPHAN INDOLE-LYASE, PYRIDOXAL 2 5-PHOSPHATE, MONOVALENT CATION BINDING SITE	TRANSFERASE TRANSFERASE, METABOLIC ROLE, PYRIDOXAL 5'- PHOSPHATE	LYASE ALPHA/BETA FOLD	TRANSFERASE TRANSFERASE, AMINOTRANSFERASE, PYRIDOXAL PHOSPHATE	TRANSFERASE SHMT; HYDROXYMETHYL TRANSFERASE, 1 CARBON METABOLISM	METHIONINE BIOSYNTHESIS BETA CYSTATHIONASE; PLP-DEPENDENT ENZYMES, METHIONINE BIOSYNTHESIS, C-S BETA 2 LYASE	LYASE CGS; LYASE, LLP-DEPENDENT ENZYMES, METHIONINE
Coumpound	RIBOSOMAL PROTEIN L37AE; CHAIN: W; RIBOSOMAL PROTEIN L37E; CHAIN: X; RIBOSOMAL PROTEIN L39E; CHAIN: Y; RIBOSOMAL PROTEIN L44E; CHAIN: Z; RIBOSOMAL PROTEIN C44E; CHAIN: Z; RIBOSOMAL PROTEIN L6; CHAIN: I;		TRYPTOPHANASE; CHAIN: A, B, C, D;	SERINE HYDROXYMETHYLTRANS FERASE; CHAIN: A;	CSDB PROTEIN; CHAIN: A;	CYSTALYSIN; CHAIN: A, B, C, D, E, F, G, H;	SERINE HYDROXYMETHYLTRANS FERASE; CHAIN: A, B;	CYSTATHIONINE BETA- LYASE; CHAIN: A, B;	CYSTATHIONINE GAMMA-SYNTHASE;
SeqFold Score									
PMF Score			-0.11	-0.06	1.00	0.00	0.00	0.16	-0.15
Verify Score			0.09	0.20	0.57	-0.09	0.25	0.25	0.33
PSI BLAST Score			1.7e-10	1.5e-67	5.1e-76	1.7e-06	3.4e-68	3.4e-43	1.7e-52
End			320	499	488	289	499	492	492
Start AA			43	27	33	61	27	58	89
Chain ID			4	A	A	A	A	A	A
PDB ID			lax4	1bj4	1c0n	1c7n	1cj0	101	lcs1
SEQ NO:			509	509	509	209	209	209	509

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			ERASI		NT	ISTER F		DOXA		METR	THES		LATE,		THEST	AMM			<u>s</u>	ron t	급.		E-5'-	IMER			, ZIZ	Z.		
		ERINE .	,r Ansfi		ENDE	R-CLL	FRIE	PYRI		ASYM	OSYN	ATE, 2	ACRY	PLE	NASC	VTE, G			THES	HYD	OFH Y	X Y	AMIN S	RIC D		VDNA	NDNA	3801		
nnotati		MT, S	HA PL YO TR		P-DEP	ULFU	NT. S	SION		ATE,	ER BI)HASC	MINO	TCO	NE BI(/HdSC			OSYN	IALDE			200	MMET		NGER	NGER			
PDB annotation	SIS	TRANSFERASE SHMT, SERINE	S, ALP	FOL CI	SE PL	RON-S	SEST	NVE	TE, 2	OFOL	CLUST	S'-PH	NE, A	ODOC	HION	5'-PH	YASE			E SEM	ASE; C	SIS, F.	, 2 PYI	, ASYI		INCF	INCE	₹- N-8 N		
	HEN	SFERA	YLASI	LIKE	SFERA	MES, I	FERA	SEC	SPHA'	HYD	FES (OXAL	YSTE	Æ-PR	MET	OXAL	ILY, L		SOPH'S	UMATI		HIL	HATE	HATE		CEX (2	LEX (2	z, UZ		
	BIOSYNTHESIS	TRAN	METHYLASE; ALPHA PLP ASPARTATE, AMINO TRANSFERASE,	(AAT)-LIKE FOLD	TRANSFERASE PLP-DEPENDENT	ENZYMES, IRON-SULFUR-CLUSTER SYNTHESIS C.S. 2 RETA I VASE	TRANSFERASE SHMT: SERINE-	GLYCINE CONVERSION, PYRIDOXAL	5'-PHOSPHATE, 2	TETRAHYDROFOLATE, ASYMMETRIC DIMER	LYASE FES CLUSTER BIOSYNTHESIS.	PYRIDOXAL 5'-PHOSPHATE,	THIOCYSTEINE, AMINOACRYLATE,	ENZYME-PRODUCT COMPLEX	LYASE METHIONINE BIOSYNTHESIS,	PYRIDOXAL 5'-PHOSPHATE, GAMMA-	2 FAMILY, LYASE		CHLOROPHYLL BIOSYNTHESIS	GLUTAMATE SEMIALDEHYDE	AMINOMOTASE; CHLOROPHYLL	BIOSYNI HESIS, PYKLIDOXAL-5-	PHOSPHATE, 2 PYRIDOXAMINE-5'-	PHOSPHATE, ASYMMETRIC DIMER		COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING PROTEIN		
	_	├ ─	2				1		_		ш	— ;					_	1						_				· <u>·</u>		
nd	 		HYDKOX YME I HYL I KANS FERASE; CHAIN: A, B, C,		AMINOTRANSFERASE;			HYDROXYMETHYLTRANS	FERASE; CHAIN: A, B, C,		L-CYSTEINE/L-CYSTINE	C-S LYASE; CHAIN: A, B;			(4)	ASE;	CHAIN: A, B, C, D, E, F, G,			70.7	AMINOI KANSFEKASE;					ER	Ä;	5	OLIGONOCLEO I IDE RINDING STTF: CHAIN: B	
Coumpound	CHAIN: A, B, C, D;	14.0	ME I E		ANSFE	B,		METH	HAIN		E/L-C	CHA;			CYSTATHIONINE	GAMMA-SYNTHASE;	В, С, Г		=	SEMIALDEHYDE	ANOFE P.	'n				QGSR ZINC FINGER	PEPTIDE; CHAIN: A;	£	OLIGONOCLEOTIDE RINDING SITE: CHAI) 1
్ర	IN: A	E	ASE; C		NOTR	CHAIN: A, B;	里	ROXY	ASE; C		STEIN	YASE			[ATHI	MA-S	ľ. A		GLUTAMATE	ALDE	2 2	CHAIN: A, B;			:	S ZINC	JDE; S	אָלָ ביי		o O
	GHA	SERINE		Ö	AMI	CHA	SERINE	HXD	FER/	ä	1-CY	C-SI			CYS	CAIM	CHA	Ħ	פונו	SEM	AMI	CHA			i	OGSI	PEPT	DUPLEA		<u>်</u>
SeqFold Score																														
PMF Score		-0.02	-		96.0		0.31				0.93				-0.05				0.00							0.17				
Verify Score		0.17			0.30		0.23			·	0.54				0.41				0.16							-0.16				
PSI BLAST Score		1.2e-77		T	1.7e-65						3.4e-47				3.4e-58			-	1.4e-10							2.7e-05				
	_	_		-			1e-65				3.4			_	3.4			-	<u></u>							2.7				
End		200		_	492		499				442				492				319							876				
Start		27			20		27				41				<u>~</u>				45							692				
Chain ID		A			⋖.		⋖				Ą				¥].	∢							А				
PDB ID		0JP1			leg5		1eji	_			lelu		_		lqgn	_			2gsa							lalh				
SEQ ID NO:		209			209		209				209				209			3	203							512				

	PDB ID	Chain ID	Start AA	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation	
ļ	Imbe		642	689	0.00085	-0.18	0.46		MYB PROTO-ONCOGENE PROTEIN; 1MBE 4	DNA BINDING PROTEIN PROTOONCOGENE PRODUCT 1MBE 12	
ļ —	lmey	U	792	876	0.00022	0.21	0.72		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	
	lath	A	691	864	1.1c-07	-0.01	0.65		QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN	
	lidy		639	694	0.0075	-0.02	0.33		MOUSE C-MYB DNA- BINDING DOMAIN REPEAT 3; CHAIN: NULL;	DNA-BINDING PROTEIN PROTOONCOGENE PRODUCT, DNA- BINDING PROTEIN	
	lmbe		642	089	0.0026	-0.36	0.22		MYB PROTO-ONCOGENE PROTEIN; 1MBE 4	DNA BINDING PROTEIN PROTOONCOGENE PRODUCT IMBE 12	
	Imse	ပ	639	694	0.0075	-0.35	0.30		COMPLEX (BINDING PROTEINDNA) C-MYB DNA-BINDING DOMAIN COMPLEXED WITH DNA IMSE 3 (NMR, MINIMIZED AVERAGE STRUCTURE) IMSE 4 IMSE 84		
	1 ubd	ပ	771	864	8.1e-06	0.16	0.04		YY1; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	

PDB annotation	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN	DNA BINDING PROTEIN PROTOONCOGENE PRODUCT IMBE 12	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN	DNA-BINDING PROTEIN PROTOONCOGENE PRODUCT, DNA- BINDING PROTEIN	DNA BINDING PROTEIN PROTOONCOGENE PRODUCT IMBE 12	
Coumpound	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B,	MYB PROTO-ONCOGENE PROTEIN; 1MBE 4	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	MOUSE C-MYB DNA- BINDING DOMAIN REPEAT 3; CHAIN; NULL;		COMPLEX (BINDING PROTEIN/DNA) C-MYB
SeqFold Score								
PMF Score	0.39	0.17	0.46	0.72	0.65	0.33	0.22	0:30
Verify Score	0.21	-0.16	-0.18	0.21	-0.01	-0.02	-0.36	-0.35
PSI BLAST Score	5.4e-06	2.7e-05	0.00085	0.00022	1.1e-07	0.0075	0.0026	0.0075
End	864	876	689	876	864	694	089	694
Start AA	692	691	642	792	692	639	642	639
Chain ID	А	¥		ပ	∀			O
PDB ID	2gli	lalh	Imbe	Imey	lalh	lidy	Imbe	Imse
SEQ NO:	512	513	513	513	513	513	513	513

PDB annotation		COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)		
Coumpound	COMPLEXED WITH DNA IMSE 3 (NMR, MINIMIZED AVERAGE STRUCTURE) IMSE 4 IMSE 84	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	TRANSFERASE(PHOSPHO TRANSFERASE) \$C-/AMP\$- DEPENDENT PROTEIN KINASE (E.C.2.7.1.37) (\$C/APLYTIC SUBUNIT) ALPHA ISOENZYME MUTANT WITH SER 139 1APM 4 REPLACED BY ALA (\\$139A\$) COMPLEX WITH THE PEPTIDE 1APM \$ INHIBITOR PKI(5-24) AND THE DETERGENT MEGA-8 1APM 6	TRANSFERASE(PHOSPHO TRANSFERASE) \$C-/AMP\$- DEPENDENT PROTEIN KINASE (E.C.2.7.1.37)
SeqFold Score				77.95	
PMF Score		0.04	0.39		0.13
Verify Score		0.16	0.21		-0.21
PSI BLAST Score		8.1e-06	5.4e-06	0	0
End		864	864	336	327
Start AA		177	769		8
Chain ID		U	A	ш	ਜ਼
PDB ID		1ubd	2gli	lapm	lapm
SEQ NO:		513	513	514	514

PDB annotation						PHOSPHOTRANSFERASE PROTEIN	PHOSPHOTRANSFERASE PROTEIN	KINASE 1CKI 18						PHOSPHOTRANSFERASE	PHOSPHOTRANSFERASE							LIPID TRANSPORT APO A-I;	LIPOPROTEIN, LIPID TRANSPORT,	CHOLESTEROL METABOLISM, 2 ATHEROSCLEROSIS, HDL, LCAT-
Coumpound	(\$C/APK\$) 1APM 3 (CATALYTIC SUBUNIT) ALPHA ISOENZYME ALITHANT WITH SED 130	MOIANI WITH SER 139 1APM 4 REPLACED BY	ALA (/S139A\$) COMPLEX WITH THE PEPTIDE 1APM	5 INHIBITOR PKI(5-24)	MEGA-8 1APM 6	CASEIN KINASE I DELTA;	CASEIN KINASE I DELTA:	ICKI 6 CHAIN: A, B; 1CKI 7	PHOSPHOTRANSFERASE	CAMP-DEPENDENT	PROIEIN KINASE	CATALYTIC SUBUNIT	ICMK 4	CASEIN KINASE-1; 1CSN 4	CASEIN KINASE-1; 1CSN 4	TRANSFERASE (PHOSPHO	TRANSFERASE) CAMP-	KINASE (F.C.2.7.1.37)	(CAPK) 1CTP 3	(CATALYTIC SUBUNIT)	1CTP 4	APOLIPOPROTEIN A-I;	CHAIN: A, B, C, D;	
SeqFold Score						285.79			75.82						293.57	77.16						69.36		
PMF Score							1.00							1.00										
Verify Score							0.64							0.73										
PSI BLAST Score						1.5e-84	1.5e-84		0				-	3.4e-78	3.4e-78	0			-			6.8e-06		
End						300	295		336					294	291	325						271		
Start AA				- 		-	3		7					1	7							69		
Chain ID						∢	4		ш							13						4		
PDB ID			-	·		lcki	1cki		lcmk					lcsn	lcsn	lctp						lavl		
SEQ NO:						514	514		514					514	514	514						520		

PDB annotation	ACTIVATION	CONTRACTILE PROTEIN TRIPLE- HELIX COILED COIL, CONTRACTILE PROTEIN	TRANSCRIPTION REGULATION SIGMA70; RNA POLYMERASE SIGMA FACTOR, TRANSCRIPTION REGULATION				LIGASE CBL, UBCH7, ZAP-70, E2,	UBIQUITIN, E3, PHOSPHORYLATION,	I BIOINTINATION PROTEIN	DEGRADATION,		LIGASE CBL, UBCH7, ZAP-70, E2,	UBIQUITIN, E3, PHOSPHORYLATION,	2 TYROSINE KINASE,	UBIQUITINATION, PROTEIN	DEGRADATION,			ZINC-BINDING PROTEIN ZINC-	BINDING PROTEIN, XNF7, BBOX,	DEVELOPMENT, 3 MID-BLASTULA-	ZINC. BINDING PROTEIN ZINC.	BINDING PROTEIN, XNF7, BBOX, DEVELOPMENT, 3 MID-BLASTULA-
Coumpound		HUMAN SKELETAL MUSCLE ALPHA-ACTININ 2; CHAIN: A;	RNA POLYMERASE PRIMARY SIGMA FACTOR; CHAIN: NULL;	VIRUS EQUINE HERPES	VIRUS-1 (C3HC4, OR RING	DOMAIN) ICHC 3 (NMK, I STRUCTURE) ICHC 4	SIGNAL TRANSDUCTION	PROTEIN CBL; CHAIN: A;	B. UBIOUTIN-	CONJUGATING ENZYME	E12-18 KDA UBCH7;	SIGNAL TRANSDUCTION	PROTEIN CBL; CHAIN: A;	ZAP-70 PEPTIDE; CHAIN:	B; UBIQUITIN-	CONJUGATING ENZYME	E12-18 KDA UBCH7;	CHAIN: C;	NUCLEAR FACTOR XNF7;	CHAIN: NULL;		NIICI FAR FACTOR XNE7.	CHAIN: NULL;
SeqFold Score		74.38	78.92																				
PMF Score				0.78			0.03					0.15							0.34			0.53	
Verify Score				0.27			-0.34					-0.19							-0.63			-0 53	
PSI BLAST Score		5.4e-12	2.2e-10	5.4e-11			1.7e-09		•			1.1e-11							5.4e-12			0.00068	
End		352	299	99			59					71							133			134	
Start		101	_	12			16					16							96			86	
Chain ID		٧					A					A											
PDB 1D		Iquu	lsig	1chc			1fbv					1fbv							1 fre			1fre	
SEQ ID NO:		520	520	526			526					526							526			526	

PDB annotation	TRANSITION	METAL BINDING PROTEIN RING FINGER PROTEIN MATI; RING IN: A: FINGER (C3HCA)	1		,; DNA-BINDING PROTEIN V(D)J RECOMBINATION ACTIVATING PROTEIN 1; RAG1, V(D)J RECOMBINATION, ANTIBODY, MAD, RING FINGER, 2 ZINC BINUCLEAR CLUSTER, ZINC FINGER, DNA- BINDING PROTEIN	OXIDE OXIDOREDUCTASE PDZ DOMAIN, A; NNOS, NITRIC OXIDE SYNTHASE AIN: B;	RIPT; PEPTIDE RECOGNITION PEPTIDE RECOGNITION, PROTEIN LOCALIZATION	HAIN: CYTOKINE LCF; CYTOKINE, LYMPHOCYTE CHEMOATTRACTANT FACTOR, PDZ DOMAIN	
Coumpound		CDK-ACTIVATING KINASE ASSEMBLY FACTOR MATICHAIN: A:	CDK-ACTIVATING KINASE ASSEMBLY FACTOR MAT1: CHAIN: A:	RAGI; CHAIN: NULL;	RAGI; CHAIN: NULL;	NEURONAL NITRIC OXIDE SYNTHASE; CHAIN: A; HEPTAPEPTIDE; CHAIN: B;	PSD-95; CHAIN: A; CRIPT; CHAIN: B;	INTERLEUKIN 16; CHAIN: NULL;	HCASK/LIN-2 PROTEIN; CHAIN: A, B;
SeqFold Score									
PMF Score		0.37	0.25	0.47	0.23	0.98	0.77	0.46	0.59
Verify Score		0.16	-0.04	-0.04	-0.28	-0.43	-0.55	-0.11	-0.52
PSI BLAST Score		1.4e-13	5.1e-05	2.4e-19	8.5e-13	1.6e-05	1.4e-05	1.1e-05	5.4e-07
End AA		7.5	99	103	101	251	251	251	260
Start		12	16	. 10	3	208	214	861	203
Chain ID		A	4			A	Ą		A
PDB ID		1g25	1g25	Irmd	lrmd	1589	1be9	1i16	lkwa
SEQ ID NO:		526	526	526	526	532	532	532	532

	Γ								(2)	
PDB annotation	KINASE	SIGNAL TRANSDUCTION HDLG, DHR3 DOMAIN; SIGNAL TRANSDUCTION, SH3 DOMAIN, REPEAT	MEMBRANE PROTEIN/OXIDOREDUCTASE BETA- FINGER, HETERODIMER	PEPTIDE RECOGNITION PSD-95; PDZ DOMAIN, NEURONAL NITRIC OXIDE SYNTHASE, NMDA RECEPTOR 2 BINDING	HYDROLASE PDZ DOMAIN, HUMAN PHOSPHATASE, HPTP1E, PTP-BAS, SPECIFICITY 2 OF BINDING	OXIDOREDUCTASE PDZ DOMAIN, NNOS, NITRIC OXIDE SYNTHASE	PEPTIDE RECOGNITION PEPTIDE RECOGNITION, PROTEIN LOCALIZATION	CYTOKINE LCF; CYTOKINE, LYMPHOCYTE CHEMOATTRACTANT FACTOR, PDZ DOMAIN	KINASE HCASK, GLGF REPEAT, DHR; PDZ DOMAIN, NEUREXIN, SYNDECAN, RECEPTOR CLUSTERING, KINASE	SIGNAL TRANSDUCTION HDLG, DHR3 DOMAIN; SIGNAL TRANSDUCTION, SH3 DOMAIN,
Coumpound		HUMAN DISCS LARGE PROTEIN; CHAIN: NULL;	ALPHA-I SYNTROPHIN (RESIDUES 77-171); CHAIN: A; NEURONAL NITRIC OXIDE SYNTHASE (RESIDUES 1-130); CHAIN:	POSTSYNAPTIC DENSITY PROTEIN 95; CHAIN: A;	TYROSINE PHOSPHATASE (PTP-BAS, TYPE 1); CHAIN: A;	NEURONAL NITRIC OXIDE SYNTHASE; CHAIN: A; HEPTAPEPTIDE; CHAIN: B;	PSD-95; CHAIN: A; CRIPT; CHAIN: B;	INTERLEUKIN 16; CHAIN: NULL;	HCASK/LIN-2 PROTEIN; CHAIN: A, B;	HUMAN DISCS LARGE PROTEIN; CHAIN: NULL;
SeqFold Score										
PMF Score		0.47	0.05	0.41	0.84	96'0	0.77	0.46	0.59	0.47
Verify Score		-0.62	-0.33	-0.38	-0.26	-0.43	-0.55	-0.11	-0.52	-0.62
PSI BLAST Score		1.6e-05	5.4e-07	5.4e-05	2.4e-05	1.6e-05	1.4e-05	1.1e-05	5.4e-07	1.6e-05
End AA		239	254	272	273	251	251	251	260	239
Start		204	199	208	208	208	214	198	203	204
Chain ID		-	∢	A	А	A	А		Ą	
PDB ID		1pdr	lqav	1qlc	3pdz	1 b 8q	1be9	1116	Ikwa	lpdr
SEQ NO:		532	532	532	532	532	532	532	532	532

Coumpound PDB annotation	REPEAT	ALPHA-1 SYNTROPHIN MEMBRANE (RESIDUES 77-171); PROTEIN/OXIDOREDUCTASE BETA-CHAIN: A; NEURONAL FINGER, HETERODIMER NITRIC OXIDE SYNTHASE (RESIDUES 1-130); CHAIN: B:	POSTSYNAPTIC DENSITY PEPTIDE RECOGNITION PSD-95; PDZ PROTEIN 95; CHAIN: A; SYNTHASE, NMDA RECEPTOR 2 BINDING	TYROSINE PHOSPHATASE HYDROLASE PDZ DOMAIN, HUMAN (PTP-BAS, TYPE I); CHAIN: PHOSPHATASE, HPTPIE, PTP-BAS, SPECIFICITY 2 OF BINDING	NEURONAL NITRIC OXIDE OXIDOREDUCTASE PDZ DOMAIN, SYNTHASE; CHAIN: A; NNOS, NITRIC OXIDE SYNTHASE HEPTAPEPTIDE; CHAIN: B:	PSD-95; CHAIN: A; CRIPT; PEPTIDE RECOGNITION PEPTIDE RECOGNITION, PROTEIN LOCALIZATION	INTERLEUKIN 16; CHAIN: CYTOKINE LCF; CYTOKINE, NULL; FACTOR, PDZ DOMAIN	2 PROTEIN;	HUMAN DISCS LARGE SIGNAL TRANSDUCTION HDLG, PROTEIN; CHAIN: NULL; TRANSDUCTION, SH3 DOMAIN, REPEAT	AT PHA. 1 CYNTTR OPHTM MEMABOANTE
		ALPHA- (RESIDU CHAIN: , NITRIC ((RESIDU B:	POSTSY	TYROSII (PTP-BA) A;	NEURON SYNTHA HEPTAPI	PSD-95; CF CHAIN: B;	INTERLE NULL;	HCASK/LIN- CHAIN: A, B;	HUMAN	ALPHA-1
SeqFold Score										
PMF Score		0.05	0.41	0.84	0.98	0.77	0.46	0.59	0.47	0.05
Verify Score		-0.33	-0.38	-0.26	-0.43	-0.55	-0.11	-0.52	-0.62	-0.33
PSI BLAST Score		5.4e-07	5.4e-05	2.4e-05	1.6e-05	1.4e-05	1.1e-05	5.4e-07	1.6e-05	5.4e-07
End AA		254	272	273	251	251	251	260	239	254
Start		199	208	208	208	214	198	203	204	199
Chain ID		¥	А	А	А	Ą		¥		Ą
PDB CI		lqav	1qlc	3pdz	168q	1be9	1116	Ikwa	1 pdr	lqav
SEQ NO:		532	532	532	533	533	533	533	533	533

PDB annotation	FINGER, HETERODIMER E :	PEPTIDE RECOGNITION PSD-95; PDZ DOMAIN, NEURONAL NITRIC OXIDE SYNTHASE, NMDA RECEPTOR 2 BINDING	HYDROLASE PDZ DOMAIN, HUMAN N: PHOSPHATASE, HPTP1E, PTP-BAS, SPECIFICITY 2 OF BINDING		PEPTIDE RECOGNITION PEPTIDE RECOGNITION, PROTEIN LOCALIZATION	 	KINASE HCASK, GLGF REPEAT, DHR; PDZ DOMAIN, NEUREXIN, SYNDECAN, RECEPTOR CLUSTERING, KINASE	SIGNAL TRANSDUCTION HDLG, DHR3 DOMAIN; SIGNAL TRANSDUCTION, SH3 DOMAIN, REPEAT	MEMBRANE PROTEIN/OXIDOREDUCTASE BETA- FINGER, HETERODIMER
Coumpound	CHAIN: A; NEURONAL NITRIC OXIDE SYNTHASE (RESIDUES 1-130); CHAIN: B;	POSTSYNAPTIC DENSITY PROTEIN 95; CHAIN: A;	TYROSINE PHOSPHATASE (PTP-BAS, TYPE 1); CHAIN: A;	NEURONAL NITRIC OXIDE SYNTHASE; CHAIN: A; HEPTAPEPTIDE; CHAIN: B:	PSD-95; CHAIN: A; CRIPT; CHAIN: B;	NTERLEUKIN 16; CHAIN: NULL;	HCASK/LIN-2 PROTEIN; CHAIN: A, B;	HUMAN DISCS LARGE PROTEIN; CHAIN: NULL;	ALPHA-1 SYNTROPHIN (RESIDUES 77-171); CHAIN: A; NEURONAL NITRIC OXIDE SYNTHASE (RESIDUES 1-130); CHAIN:
SeqFold Score									
PMF Score		0.41	0.84	0.98	0.77	0.46	0.59	0.47	0.05
Verify Score		-0.38	-0.26	-0.43	-0.55	-0.11	-0.52	-0.62	-0.33
PSI BLAST Score		5.4e-05	2.4e-05	1.6e-05	1.4e-05	1.1e-05	5.4e-07	1.6e-05	5.4e-07
End		272	273	251	251	251	260	239	254
Start		208	208	208	214	861	203	204	199
Chain ID		Ą	А	¥	¥		Ą		A
PDB ID		1qlc	3pdz	158գ	1be9	1116	1kwa	1pdr	Iqav
SEQ ID NO:		533	533	533	533	533	533	533	533

SEQ B D	PDB ID	Chain ID	Start	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
.	1qlc	∢	208	272	5.4e-05	-0.38	0.41		POSTSYNAPTIC DENSITY PROTEIN 95; CHAIN: A;	PEPTIDE RECOGNITION PSD-95; PDZ DOMAIN, NEURONAL NITRIC OXIDE SYNTHASE, NMDA RECEPTOR 2 RINDING
533	3pdz	4	208	273	2.4e-05	-0.26	0.84		TYROSINE PHOSPHATASE (PTP-BAS, TYPE 1); CHAIN: A;	HYDROLASE PDZ DOMAIN, HUMAN PHOSPHATASE, HPTP1E, PTP-BAS, SPECIFICITY 2 OF BINDING
538	1c4z	A	234	583	0	0.33	1.00		UBIQUITIN-PROTEIN LIGASE E3A; CHAIN: A, B, C; UBIQUITIN CONJUGATING ENZYME	LIGASE E6AP; UBCH7; BILOBAL STRUCTURE, ELONGATED SHAPE, E3 UBIQUITIN LIGASE, E2 2 UBIQUITIN CONJUGATING ENZYME
538	1c4z	⋖	234	586	0			227.88	UBIQUITIN-PROTEIN LIGASE E34; CHAIN: A, B, C; UBIQUITIN CONJUGATING ENZYME E2; CHAIN: D;	LIGASE E6AP; UBCH7; BILOBAL STRUCTURE, ELONGATED SHAPE, E3 UBIQUITIN LIGASE, E2 2 UBIQUITIN CONJUGATING ENZYME
	la7i		100	159	1.4e-15	69:0	0.94		QCRP2 (LIM1); CHAIN: NULL;	LIM DOMAIN CONTAINING PROTEINS LIM DOMAIN CONTAINING PROTEINS, METAL-BINDING PROTEIN ZINC 2 FINGER
	la7i		163	218	1.9e-13	-0.03	0.83		QCRP2 (LIM1); CHAIN: NULL;	LIM DOMAIN CONTAINING PROTEINS LIM DOMAIN CONTAINING PROTEINS, METAL-BINDING PROTEIN, ZINC 2 FINGER
	la7i		39	94	5.4e-14	0.40	0.93		QCRP2 (LIM1); CHAIN: NULL;	LIM DOMAIN CONTAINING PROTEINS LIM DOMAIN CONTAINING PROTEINS, METAL-BINDING PROTEIN, ZINC 2 FINGER
	1b8t	A	162	281	3.4e-13	0.01	0.84		CRPI; CHAIN: A;	CONTRACTILE LIM DOMAIN, CRP, NMR, MUSCLE DIFFERENTIATION, CONTRACTILE

		Start	End	PSI RLAST	Verify	PMF	SeqFold	Coumpound	PDB annotation
-	3		1	Score	3	1000			
	A	33	236	5.1e-14			60.96	CRPI; CHAIN: A;	CONTRACTILE LIM DOMAIN, CRP, NMR, MUSCLE DIFFERENTIATION, CONTRACTILE
		38	25	5.4e-16	0.19	1.00		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS ICTL 15
<u> </u>		66	155	5.4e-13	0.30	0.82		AVIAN CYSTEINE RICH PROTEIN; ICTL 3	METAL-BINDING PROTEIN LIM DOMAIN CONTAINING PROTEINS 1CTL 15
	A	160	215	2.7e-13	0.44	0.71		CYSTEINE AND GLYCINE- RICH PROTEIN CRP2; CHAIN: A;	SIGNALING PROTEIN LIM DOMAIN CONTAINING PROTEINS, METAL- BINDING PROTEIN
	A	38	94	1.6e-15	0.74	1.00		CYSTEINE AND GLYCINE- RICH PROTEIN CRP2; CHAIN: A;	SIGNALING PROTEIN LIM DOMAIN CONTAINING PROTEINS, METAL- BINDING PROTEIN
	A	66	157	1.4e-14	0.10	86.0		CYSTEINE AND GLYCINE- RICH PROTEIN CRP2; CHAIN: A;	SIGNALING PROTEIN LIM DOMAIN CONTAINING PROTEINS, METAL- BINDING PROTEIN
	∢	104	270	6.8e-07			59.59	TUMOR NECROSIS FACTOR RECEPTOR; CHAIN: A, B;	SIGNALLING PROTEIN BINDING PROTEIN, CYTOKINE, SIGNALLING PROTEIN
		101	170	1.le-15	0.12	0.41		CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NULL;	METAL-BINDING PROTEIN CRIP; METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN
		163	230	5.4e-21	0.05	0.64		CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NULL;	METAL-BINDING PROTEIN CRIP; METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN
		41	111	5.4e-17	0.27	0.27		CYSTEINE RICH INTESTINAL PROTEIN; CHAIN: NULL;	METAL-BINDING PROTEIN CRIP; METAL-BINDING PROTEIN, LIM DOMAIN PROTEIN
\vdash		72	241	0.00027			62.78	LAMININ; CHAIN: NULL;	GLYCOPROTEIN GLYCOPROTEIN
	A		991	3.4e-07			52.60	TUMOR NECROSIS FACTOR RECEPTOR; INCF 4 CHAIN: A. B: INCF 5	SIGNALLING PROTEIN TYPE I RECEPTOR, STNFRI; INCF 8 BINDING PROTEIN CYTOKINE INCF 19

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
542	labr	Я	529	929	5.1e-13	-0.43	0.27		COMPLEX (GLYCOSIDASE/CARBOHY DRATE) ABRIN-A COMPLEXED WITH TWO SUGAR CHAINS 1ABR 3	
542	lqgq	Y	191	431	1.4e-21	0.13	0.58		SPORE COAT POLYSACCHARIDE BIOSYNTHESIS PROTEIN CHAIN: A;	TRANSFERASE GLYCOSYLTRANSFERASE
542	lxyf	А	496	629	1.7e-30	0.30	69.0		ENDO-1,4-BETA- XYLANASE; CHAIN: A, B;	HYDROLASE XYLAN DEGRADATION
542	lxyf	A	546	969	16-17	0.04	0.70		ENDO-1,4-BETA- XYLANASE; CHAIN: A, B;	HYDROLASE XYLAN DEGRADATION
543	1c4o	A	559	681	5.4e-11	0.25	0.65		DNA NUCLEOTIDE EXCISION REPAIR ENZYME UVRB; CHAIN: A;	REPLICATION DNA NUCLEOTIDE EXCISION REPAIR, UVRABC, HELICASE, 2 HYPERTHERMOSTABLE PROTEIN
543	1d2m	A	559	681	2.7e-10	0.23	0.54		EXCINUCLEASE ABC SUBUNIT B; CHAIN: A;	HYDROLASE UVRB; MULTIDOMAIN PROTEIN
543	149x	A	536	683	2.4e-18	0.61	1.00		EXCINUCLEASE UVRABC COMPONENT UVRB; CHAIN: A;	GENE REGULATION APO PROTEIN
543	1fuk	Ą	544	682	1.1e-11	0.71	09.0		EUKARYOTIC INITIATION FACTOR 4A; CHAIN: A;	TRANSLATION YEAST INITIATION FACTOR 4A, EIF4A; HELICASE, INITIATION FACTOR 4A, DEAD-BOX PROTEIN
543	1fuu	æ	555	652	1.6e-11	0.36	0.62		YEAST INITIATION FACTOR 4A; CHAIN: A, B;	TRANSLATION EUKARYOTIC INITIATION FACTOR 44; IF4A, HELICASE, DEAD-BOX PROTEIN
546	le0g	∢	65	107	8.5e-08	-0.72	0.04		MEMBRANE-BOUND LYTIC MUREIN	HYDROLASE MLTD, MUREIN HYDROLASE D, REGULATORY

PDB annotation	PROTEIN DNIR; CELL WALL, HYDROLASE, GLYCOSIDASE, LIPOPROTEIN, 2 OUTER MEMBRANE, MULTIGENE FAMILY		COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN		COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA	INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	MITTER A CITION DE CITE DE SIGN 2	CRYSTAL STRUCTURE, COMPLEX	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	COVERACTION, PROTEIN DESIGN, 2	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	Chycara capitoring commy	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2
Coumpound	TRANSGLYCOSYLASE D; CHAIN: A;	i	QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE RINDING SITE: CHAIN: B	C. C.	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	PROTEIN: CHAIN: OF G.	A MOTERIA, CLEMIN. C, I., C,		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	raciein, chain. C. F. G.		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;
SeqFold Score						•				•		-									
PMF Score			0.86		1.00			1.00	-			1.00				66.0	-			69.0	
Verify Score			-0.04		0.16			0.32				0.16				0.61				0.02	
PSI BLAST Score			1.4e-27		8.5e-44			3.4e-47		•		8.5e-48				1.7e-46	•			1.1e-39	
End			351		182			238				566				293				351	
Start AA			270		101			157				185				213				241	
Chain ID			V		O			ပ				ပ				ပ				ပ	
PDB ID			laih		lmey			lmey				Imey				Imey				lmey	
SEQ ID NO:			250		550			220				250				520				220	

SEQ	PDB	п	•	End	PSI DI ACT	Verify	PMF	SeqFold	Coumpound	PDB annotation
	a	9	AA	AA	Score	Score	Score	Score		
				,						CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)
550	lmey	၁	569	351	1.7e-46	-0.06	1.00		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA
									PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX
Π.										(ZINC FINGER/DNA)
250	lmey	ပ	298	379	1.5e-49	0.16	1.00	,	DNA; CHAIN: A, B, D, E; CONSENSIIS ZINC FINGER	COMPLEX (ZINC FINGER/DNA) ZINC FINGER PROTEIN, DNA
						_			PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
										CRYSTAL STRUCTURE, COMPLEX
550	Imey	ပ	326	407	1.4e-50	0.29	1.00		DNA CHAIN A B D E	COMPLEX (ZINC FINGER (DNA) ZINC
			·						CONSENSUS ZINC FINGER	FINGER, PROTEIN-DNA
									PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
										CRYSTAL STRUCTURE, COMPLEX
1		ļ								(ZINC FINGER/DNA)
کر —	Imey	ပ	354	435	8.5e-51	0.21	0.0		DNA; CHAIN: A, B, D, E;	COMPLEX (ZINC FINGER/DNA) ZINC
									CONSENSUS ZINC FINGER	FINGER, PROTEIN-DNA
_						_			PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
_								•		CRYSTAL STRUCTURE, COMPLEX
7										(ZINC FINGER/DNA)
220	Imey	ပ_	382	63	1e-50	0.21	1.00		DNA; CHAIN: A, B, D, E;	COMPLEX (ZINC FINGER/DNA) ZINC
					-				CONSENSUS ZINC FINGER	FINGER, PROTEIN-DNA
_			_		•				PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
			_			_				CRYSTAL STRUCTURE, COMPLEX
7			1							(ZINC FINGER/DNA)
550	Imey	ပ	410	491	1.7e-50	0.16	1.00		DNA; CHAIN: A, B, D, E;	COMPLEX (ZINC FINGER/DNA) ZINC
_									CONSENSUS ZINC FINGER	FINGER, PROTEIN-DNA
_		_							PROTEIN; CHAIN: C, F, G;	INTERACTION, PROTEIN DESIGN, 2
										CRYSTAL STRUCTURE, COMPLEX
†		 	1	1						(ZINC FINGER/DNA)
000	Imey	۔۔۔۔۔	438	519	le-50	0.29	 8:		DNA; CHAIN: A, B, D, E;	COMPLEX (ZINC FINGER/DNA) ZINC FINGER PROTEIN, DNA
1			1	1					CONDENSOD FINO I INOUN	INGER, I ROLDIN-DIAR

PDB annotation	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DINA)		INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER PROTFIN-DNA		CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC		INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC		INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	COMPLEY (TO ANSCRIPTION	REGULATION/DNA) COMPLEX:	(TRANSCRIPTION	REGULATION/DNA), RNA POI VACED ASE III 3 TRANSCRIPTION
Coumpound	PROTEIN; CHAIN: C, F, G;	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		TFIIIA; CHAIN: A, D; 5S	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;			TEITIA: CHAIN: A D: 50	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;	
SeqFold Score		107.02																			107.82	30.701		
PMF Score					1.00			1.00				0.65				0.59								
Verify Score					0.29			0.20				-0.44				-0.13								
PSI BLAST Score		1e-50			5.1e-50			8.5e-35				1e-33				5.1e-35					3 40-36	3		
End		520			547			552				154				247					205	}		
Start AA		438			466			464				85				102					127	į		
Chain ID		၁			ပ			ပ				ပ				A					A	;		
PDB ID		lmey			Imey			lmey				Imey				1tf6					1166			
SEQ ID NO:		250			550			550				550				220					250	2		

PDB ET	Chain ID	Start AA	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
- 1				Score					TOTAL AND THE PARTY OF THE PARTY AND THE PAR
1	A	158	304	3.4e-34	0.18	96.0		TFIIIA; CHAIN: A, D; 5S	COMPLEX (TRANSCRIPTION
								KIBOSOMAL KNA GENE; CHAIN: B, C, E, F;	KEGULA HON/DNA) COMPLEX (TRANSCRIPTION
									REGULATION/DNA), RNA
- 1									POLYMERASE III, 2 TRANSCRIPTION INITIATION, ZINC FINGER PROTEIN
1tf6	A	186	332	1.7e-34	-0.21	0.70		TFIIIA; CHAIN: A, D; 5S	COMPLEX (TRANSCRIPTION
								RIBOSOMAL RNA GENE; CHAIN: B. C. E. F:	REGULATION/DNA) COMPLEX (TRANSCRIPTION
									REGULATION/DNA), RNA
									POLYMERASE III, 2 TRANSCRIPTION
11f6	A	383	529	le-37	0.05	96.0		TFIIIA: CHAIN: A. D: 5S	COMPLEX (TRANSCRIPTION
								RIBOSOMAL RNA GENE;	REGULATION/DNA) COMPLEX
								CHAIN: B, C, E, F;	(TRANSCRIPTION
									REGULATION/DNA), RNA
									POLYMERASE III, 2 TRANSCRIPTION
I.	_		3	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	100	;			INITIATION, CINC FINGER PROTEIN
911	∢	114	249	3.4e-36	0.21	0.84		TFIIIA; CHAIN: A, D; 5S RIBOSOMAL RNA GENE;	COMPLEX (TRANSCRIPTION REGULATION/DNA) COMPLEX
					,,			CHAIN: B, C, E, F;	(TRANSCRIPTION
									REGULATION/DNA), RNA
							-		POLYMERASE III, 2 TRANSCRIPTION
1tf6	A	93	219	1.7e-30	-0.00	0.89		TFIIIA; CHAIN: A, D; 5S	COMPLEX (TRANSCRIPTION
								RIBOSOMAL RNA GENE;	REGULATION/DNA) COMPLEX
								CHAIN: B, C, E, F;	(TRANSCRIPTION
									REGULATION/DNA), RNA
									POLYMERASE III, 2 TRANSCRIPTION
Ι.	,	1	;						INITIATION, ZINC FINGER PROTEIN
pqnI	<u>ာ</u>	104	210	6.8e-32	0.17	0.0		YY1; CHAIN: C; ADENO-	COMPLEX (TRANSCRIPTION
			·					ASSOCIATED VIRUS PS	REGULATION/DNA) YING-YANG 1;
			1					INTITUTION PROMINI	I RAINSCRIPTION INTITATION,

				~		
PDB annotation	INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATIONDNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGIL ATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATIONDNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGIT ATTONDNA)	COMPLEX (TRANSCRIPTION REGULATIONDA) YING-YANG 1;
Coumpound	DNA; CHAIN: A, B;	YY I; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY1; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS PS INITIATOR ELEMENT DNA; CHAIN: A, B;	YY I; CHAIN: C; ADENO-ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5
SeqFold Score						
PMF Score		0.90	1.00	66.0	0.75	96.0
Verify Score		-0.08	-0.21	0.27	-0.20	0.11
PSI BLAST Score		8.1e-45	2.7e-54	5.4e-56	5.4e-52	6.8e-31
End		210	238	266	292	323
Start AA		II .	134	155	183	221
Chain ID		O	U	ပ	ပ	ပ
PDB ID		lubd	lubd	1ubd	1ubd	lubd
SEQ ID NO:		550	550	550	550	550

PDB annotation	TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATIONDNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION
Coumpound	INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY I; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY1; CHAIN: C; ADENO-ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY1; CHAIN: C; ADENO-
SeqFold Score						90.51
PMF Score		0.17	0.81	0.99	1.00	
Verify Score		-0.31	-0.16	0.23	0.14	
PSI BLAST Score		5.4e-50	1.4e-33	1.16-51	1.46-58	1.4e-58
End AA		380	379	407	491	520
Start AA		239	277	596	381	410
Chain ID		ပ	ပ	ပ	ပ	ပ
PDB ID		lubd	lubd	lubd	lubd	lubd
SEQ No.		550	550	550	550	550

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PDB annotation	REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	TRANSCRIPTION REGULATION TRANSCRIPTION REGULATION, ADR1, ZINC FINGER, NMR	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLJ; GLJ, ZINC FINGER, COMPLEX (DNA-
Coumpound	ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY1; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY1; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	ADR1; CHAIN: NULL;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;
SeqFold Score							100.42
PMF Score		0.99	0.99	0.65	06'0	1.00	
Verify Score		0.04	0.12	0.08	0.09	0.37	
PSI BLAST Score		8.1e-56	6.8e-35	8.5e-16	2.7e-43	1.1e-70	1.1e-70
End AA		547	547	297	212	268	296
Start AA		436	446	242	111	130	157
Chain ID		ပ	ပ		A	А	Ą
PDB ID		lubd	lubd	2adr	2gli	2gli	2gli
SEQ NO:		550	250	550	250	550	550

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PDB annotation	BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING	ZINC FINGER, COMPLEX (DNA-	SINDING PROTEIN/DINA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI,	ZINC FINGER, COMPLEX (DNA-	BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING	PROTEIN/DNA) FIVE-FINGER GLI; GLI,	LINC FINGER, COMPLEX (DNA-	COMPLEX (DNA-BINDING	PROTEIN/DNA) FIVE-FINGER GLI: GLI.	ZINC FINGER, COMPLEX (DNA-	BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING	PROTEIN/DNA) FIVE-FINGER GLI; GLI,	ZINC FINGER, COMPLEX (DNA-	BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING	PROTEIN/DNA) FIVE-FINGER GLJ; GLJ,	ZINC FINGER, COMPLEX (DNA-		SIGNALING PROTEIN REGULATION	GALPHA INTERACTING PROTEIN;	GAIP, RGS, REGULATOR OF G	PROTEIN, SIGNALING PROTEIN 2	REGULATION	SIGNALING PROTEIN REGULATION	GALPHA INTERACTING PROTEIN;
Coumpound		ZINC FINGER PROTEIN	CHAIN: C, D;	ZNIC ENICEB BROTERI	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA;	CHAIN: C, D;		ZINC FINGER PROTEIN	GLII; CHAIN: A; DNA;	Chain: C, D;	ZINC FINGER PROTEIN	GLII; CHAIN: A; DNA;	CHAIN: C, D;		ZINC FINGER PROTEIN	GLII; CHAIN: A; DNA;	CHAIN: C, D;		ZINC FINGER PROTEIN	GLII; CHAIN: A; DNA;	CHAIN: C, D;		GAIP (G-ALPHA	INTERACTING) PROTEIN;	CHAIN: A;			GAIP (G-ALPHA	INTERACTING) PROTEIN;
SeqFold Score																									_		\dashv	172.06	-
PMF Score		1.00		370	0.40			0.58			1.00				0.88				0.99				1.00						
Verify Score		0.40		0.10	0.10			0.18			0.39				-0.09				0.17				0.47						
PSI BLAST Score		2.2e-67	-	160.65	1.06-03	-		8.5e-33			2.7e-68				8.1e-73			\neg	6.8e-35				1.4e-51					1.4e-51	
End AA		297		252			į	378			437				549				518				200					200	
Start AA		157		185	<u>-</u>		3,0	249			298				382				330	-			73					23	
Chain ID		A			<			∢			A				٧				٧				A					∀	
PDB ID		2gli		2oli	2,E11			ıı87			2gli				2gli				2gli				lcmz					lcmz	
SEQ NO:		550		550	3		3	066			550				550				220				553					523	

	2	TON N;	Z Ž	7	ESIS	Z Z	E2		ESIS	1	NPF	2 2	EIN,		بدا	SE SE	MAL			ING.	
PDB annotation	ALING PROTEIN	ATEIN REGULAT ACTING PROTEI ULATOR OF G ALING PROTEIN	BINDING PROTE	JEGI MEMBRAN PHOLIPID I GI_DERIYED	ESICLE BIOGEN	BINDING PROTI	LGI MEMBRAN	HOLIPID	ESICLE BIOGEN	TEIN CALCIUM	LING DOMAIN,	NDING, 2 EF-HA	ROTEIN G-PROT		A, GTPASE, RNA	IKE, HYDROLA!	NG EH2, EPIDEF	R RECEPTOR	CIUM BINDING	SIGNALING DOMAIN, NPF BINDING,	
PDB	PROTEIN, SIGNALING PROTEIN 2 REGULATION	SIGNALING PROTEIN REGULATION GALPHA INTERACTING PROTEIN; GAIP, RGS, REGULATOR OF G PROTEIN, SIGNALING PROTEIN 2 REGULATION	PHOSPHOLIPID-BINDING PROTEIN, PHOSPHOLIPID-BINDING PROTEIN,	PROTEIN, PHOSPHOLIPID FYCHANGE GOI GLINERIVED	SECRETORY 3 VESICLE BIOGENESIS	PHOSPHOLIPID-BINDING PROTEIN PHOSPHOLIPID-BINDING PROTEIN,	PERIPHERAL GOLGI MEMBRANE 2	PROTEIN, PHOSPHOLIPID	SECRETORY 3 VESICLE BIOGENESIS	SIGNALING PROTEIN CALCIUM	BINDING, SIGNALING DOMAIN, NPF	BINDING, FW BINDING, 2 EF-HAND, FH DOMAIN SIGNAI ING PROTEIN	RNA BINDING PROTEIN G-PROTEIN,	BETA-BARREL	HYDROLASE ERA, GTPASE, RNA-	BINDING, RAS-LIKE, HYDROLASE	CALCIUM BINDING EH2, EPIDERMAL	GROWTH FACTOR RECEPTOR	SUBSTRATE CALCIUM BINDING,	SIGNALING DON	
Coumpound		GAIP (G-ALPHA INTERACTING) PROTEIN; CHAIN: A;	PHOSPHATIDYLINOSITOL TRANSFER PROTEIN	SECTOR CIRCLES		PHOSPHATIDYLINOSITOL TRANSFER PROTEIN	SEC14P; CHAIN: NULL;			EPIDERMAL GROWTH	FACTOR RECEPTOR	FAIRWAY CHAIN; A;	ELONGATION FACTOR TU	(EF-TU); CHAIN: A, B, C, D	GTP-BINDING PROTEIN	ERA; CHAIN: A, B;	EPS15; CHAIN; NULL;				
SeqFold Score			92.44													i	94.67		-	_	_
PMF Score		1.00				1.00				1.00			0.03		0.09					_	
Verify Score		0.47				0.22				0.87			-0.44		-0.24					_	
PSI BLAST Score		3.4e-47	8.5e-61			8.5e-61				1.7e-19			1.4e-83		0.001		1.7e-39				
End AA		200	277			274				534			424		263		536			_	
Start AA		73	17			43				444 			56		61		443				
Chain ID		Ą								∢			A		Ą						
PDB ID		1cmz	laua			lana				1007			1d2e	,	lega		leh2				
SEQ NO:		553	555			555				557			557		557		557				

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PDB annotation	CALCIUM BINDING EH2, EPIDERMAL GROWTH FACTOR RECEPTOR SUBSTRATE CALCIUM BINDING, SIGNALING DOMAIN, NPF BINDING, EF-HAND, EH 2 DOMAIN		TRANSLATION TRANSLATIONAL GTPASE		PROTEIN BINDING EF-G; EF-G ELONGATION FACTOR, TRANSLOCASE, RIBOSOME, ELONGATION, 2 TRANSLATION, PROTEIN SYNT FACTOR, GTPASE, GTP BINDING, 3 GUANOSINE NUCLEOTIDE BINDING, PROTEIN BINDING	RIBOSOME 50S RIBOSOMAL PROTEIN L2P, HMAL2, HL4; 50S RIBOSOMAL PROTEIN L3P, HMAL3, HL1; 50S RIBOSOMAL PROTEIN L4E, HMAL4, HL6; 50S RIBOSOMAL PROTEIN L5P, HMAL5, HL13; 30S RIBOSOMAL PROTEIN HS6; 50S RIBOSOMAL
Coumpound	EPSIS; CHAIN: NULL;	TRANSPORT AND PROTECTION PROTEIN ELONGATION FACTOR TU (DOMAIN I) - *GUANOSINE DIPHOSPHATE IETU 4 COMPLEX IETU 5	TRANSLATION INITIATION FACTOR IF2/EIF5B; CHAIN: A;	CALCIUM-BINDING PROTEIN RAT ONCOMODULIN IRRO 3	ELONGATION FACTOR G; CHAIN: A; ELONGATION FACTOR G DOMAIN 3; CHAIN: B;	23S RRNA; CHAIN: 0; 5S RRNA; CHAIN: 9; RIBOSOMAL PROTEIN L2; CHAIN: A; RIBOSOMAL PROTEIN L3; CHAIN: B; RIBOSOMAL PROTEIN L4; CHAIN: C; RIBOSOMAL
SeqFold Score						
PMF Score	1.00	0.11	0.05	0.13	0.10	0.18
Verify Score	1.11	-0.02	-0.05	0.06	-0.40	-0.34
PSI BLAST Score	1.7e-39	1e-52	6.8e-13	0.0011	1.4e-13	5.1e-30
End	536	284	331	506	222	92
Start	444	52	65	428	56	2
Chain ID			A		V	2
PDB ID	leh2	letu	1g7s	Irro	2efg	IŘ
SEQ ID NO:	557	557	557	557	557	559

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PDB annotation	PROTEIN L13P, HMAL13; 50S RIBOSOMAL PROTEIN L14P, HMAL14,	HL27; 50S RIBOSOMAL PROTEIN L15P,	HMAL15, HL9; 50S RIBOSOMAL	PROTEIN LISP, HMALIS, HL12; 50S	RIBOSOMAL PROTEIN L18E, HL29,	L19; 50S RIBOSOMAL PROTEIN L19E,	HMAL19, HL24; 50S RIBOSOMAL	PROTEIN L21E, HL31; 50S RIBOSOMAL	FROIEIN LZZP, HMALZZ, HLZZ; 50S	III 26 121 605 BID COSE, HMALZS,	LLCS, LCI; 30S KIBOSOWAL PROTEIN	PIBOSOMAI DROTTENI 24E	HI 21/HI 22: 500 PIBOSOMAI BROTEIN	L29P. HMAL29, HL33: 50S RIBOSOMAL	PROTEIN L30P, HMAL30, HL20, HL16;	50S RIBOSOMAL PROTEIN L31E, L34,	HL30; 50S RIBOSOMAL PROTEIN L32E,	HL5; 50S RIBOSOMAL PROTEIN L37E,	L35E; 50S RIBOSOMAL PROTEINS	L39E, HL39E, HL46E; 50S RIBOSOMAL	PROTEIN L44E, LA, HLA; 50S	RIBOSOMAL PROTEIN L6P, HMAL6,	HLIO RIBOSOME ASSEMBLY, RNA.	KNA, PROTEIN-RNA, PROTEIN-	PROTEIN							
Coumpound	PROTEIN L5; CHAIN: D; RIBOSOMAL PROTEIN	L7AE; CHAIN: E;	RIBOSOMAL PROTEIN	L10E; CHAIN: F;	KIBOSOMAL PROTEIN	L13; CHAIN: G;	RIBOSOMAL PROTEIN	L14; CHAIN: H;	KIBOSOMAL PROJEIN	LIJE; CHAIN: I;	NEOSOWAL PROTEIN	RIBOSOMAI PROTEIN	T 18: CHAIN: K.	RIBOSOMAL PROTEIN	L18E; CHAIN: L;	RIBOSOMAL PROTEIN	L19; CHAIN: M;	RIBOSOMAL PROTEIN	L21E; CHAIN: N;	RIBOSOMAL PROTEIN	L22; CHAIN: 0;	RIBOSOMAL PROTEIN	L23; CHAIN: P;	KIBOSOMAL PROTEIN	L24; CHAIN: Q;	RIBOSOMAL PROTEIN	L24E; CHAIN: R;	RIBOSOMAL PROTEIN	L29; CHAIN: S;	RIBOSOMAL PROTEIN	L30; CHAIN: 1;	KIBOSOMAL PROTEIN
SeqFold Score																											_					
PMF Score																													•			
Verify Score																																
PSI BLAST Score																						-										
End AA																		·														
Start AA																																
Chain ID																																
PDB CI																		-														
SEQ NO:															_																	

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		Τ								_			_	_					_															_
PDB annotation								•						RIBOSOME 50S RIBOSOMAL PROTEIN	L2P, HMAL2, HL4; 50S RIBOSOMAL	PROTEIN L3P, HMAL3, HL1; 50S	RIBOSOMAL PROTEIN L4E, HMAL4,	HL6; 50S RIBOSOMAL PROTEIN LSP,	HMAL5, HL13; 30S RIBOSOMAL	PROTEIN HS6; 50S RIBOSOMAL	PROTEIN L13P, HMAL13; 50S	RIBOSOMAL PROTEIN L14P, HMAL14,	HL27; 50S RIBOSOMAL PROTEIN L15P,	HMAL15, HL9; 50S RIBOSOMAL	PROTEIN L18P, HMAL18, HL12; 50S	RIBOSOMAL PROTEIN L18E, HL29,	L19; 50S RIBOSOMAL PROTEIN L19E,	HMAL19, HL24; 50S RIBOSOMAL	PROTEIN L21E, HL31; 50S RIBOSOMAL	PROTEIN L22P, HMAL22, HL23; 50S	RIBOSOMAL PROTEIN L23P, HMAL23,	HL25, L21; 50S RIBOSOMAL PROTEIN	L24P, HMAL24, HL16, HL15; 50S	TATOSOMAT DE OTTETATIONE
Coumpound		L31E; CHAIN: U; RIBOSOMAL PROTEIN	L32E; CHAIN: V;	RIBOSOMAL PROTEIN	L37AE; CHAIN: W;	RIBOSOMAL PROTEIN	L37E; CHAIN: X;	RIBOSOMAL PROTEIN	L39E; CHAIN: Y;	RIBOSOMAL PROTEIN	L44E; CHAIN: Z;	RIBOSOMAL PROTEIN L6;	CHAIN: 1;	23S RRNA; CHAIN: 0; 5S	RRNA; CHAIN: 9;	RIBOSOMAL PROTEIN L2;	CHAIN: A; RIBOSOMAL	PROTEIN L3; CHAIN: B;	RIBOSOMAL PROTEIN L4;	CHAIN: C; RIBOSOMAL	PROTEIN L5; CHAIN: D;	RIBOSOMAL PROTEIN	L7AE; CHAIN: E;	RIBOSOMAL PROTEIN	L10E; CHAIN: F;	RIBOSOMAL PROTEIN	L13; CHAIN: G;	RIBOSOMAL PROTEIN	L14; CHAIN: H;	RIBOSOMAL PROTEIN	L15E; CHAIN: I;	RIBOSOMAL PROTEIN	L15; CHAIN: J;	PIROCOMAT DECIMA
SeqFold	2006						-					_											-											-
PMF	21020													0.03									-											-
Verify	Score													-0.53																				
PSI RI AST	Score													2.7e-45							_		•											_
End														23													-		_					
Start														4																				
Chain	}													2																				_
PDB EDB														1#K														_						
SEQ ID	Ö													529																	,			

PDB annotation		HL21/HL22; 50S RIBOSOMAL PROTEIN L29P, HMAL29, HL33; 50S RIBOSOMAL	PROTEIN L30P, HMAL30, HL20, HL16;	50S RIBOSOMAL PROTEIN L31E, L34,	HL30; 50S RIBOSOMAL PROTEIN L32E,	HLS; 50S RIBOSOMAL PROTEIN L37E,	L35E; 50S RIBOSOMAL PROTEINS	L39E, HL39E, HL46E; 50S RIBOSOMAL	PROTEIN L44E, LA, HLA; 50S	RIBOSOMAL PROTEIN L6P, HMAL6,	HL10 RIBOSOME ASSEMBLY, RNA-	RNA, PROTEIN-RNA, PROTEIN-	PROTEIN							-												
Coumpound		LI8; CHAIN: K; RIBOSOMAL PROTEIN	L18E; CHAIN: L;	RIBOSOMAL PROTEIN	L19; CHAIN: M;	RIBOSOMAL PROTEIN	L21E; CHAIN: N;	RIBOSOMAL PROTEIN	L22; CHAIN: 0;	RIBOSOMAL PROTEIN	L23; CHAIN: P;	RIBOSOMAL PROTEIN	L24; CHAIN: Q;	RIBOSOMAL PROTEIN	L24E; CHAIN: R;	RIBOSOMAL PROTEIN	L29; CHAIN: S;	RIBOSOMAL PROTEIN	L30; CHAIN: T;	RIBOSOMAL PROTEIN	L31E; CHAIN: U;	RIBOSOMAL PROTEIN	L32E; CHAIN: V;	RIBOSOMAL PROTEIN	L37AE; CHAIN: W;	KIBOSOMAL PROTEIN	PIBOCOMAI DECTERI	NECONIAL FROIDIN	EJJE, CHAIN. I., RIBOSOMAL PROTEIN	L44E: CHAIN: Z:	RIBOSOMAL PROTEIN L6; CHAIN: 1;	
SeqFold	Score												_,_									-										
PMF	Score																								-							
Verify	Score																			··-												
PSI	BLAST Score																												<u></u> -			
End	ΑA																							•		-	_					
Start	AA						•							•																	-	
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PDB	a																							_								
SEQ	NO.																															

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PDB annotation	SIGNALING PROTEIN BETA-ALPHA- BETA FOLD PARALLEL BETA SHEET	SIGNALING PROTEIN BETA-ALPHA- BETA FOLD		IMMUNE SYSTEM IMMUNOGLOBULIN; IMMUNOGLOBULIN ANTIBODY ENGINEERING, HUMANIZED AND CHIMERIC ANTIBODY, FAB, 2 X-RAY STRUCTURE, THREE-DIMENSIONAL STRYCTURE, GAMMA- 3 NTERFERON IMMINE SYSTEM	MAINOGI OBIII IN	IMMUNOGLOBULIN, KAPPA LIGHT- CHAIN DIMER HEADER	COMPLEX (MHC/VIRAL	PEPTIDE/RECEPTOR) HLA A2 HEAVY CHAIN; COMPLEX (MHC/VIRAL	PEPTIDE/RECEPTOR)			COMPLEX (ANTIBODY/ANTIGEN)	(ANTIBODY/ANTIGEN), ANGIOGENIC	FACTOR	IMMUNE SYSTEM ANTIBODY (FAB	FKAGMEN 1), IMMONE SYSTEM		IMMUNE SYSTEM FAB-IBP COMPLEX CRYSTAL STRUCTURE 2.7A
Coumpound	TOLL-LIKE RECEPTOR 1; CHAIN: A;	TOLL-LIKE RECEPTOR 2; CHAIN: A;		ANTIBODY (LIGHT CHAIN), CHAIN: L; ANTIBODY (HEAVY CHAIN); CHAIN: H;	IMMI INOGI OBI II IN:	CHAIN: A, B;	HLA-A 0201: CHAIN: A:	BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE;	CHAIN: C; T CELL RECEPTOR ALPHA:	CHAIN: D; T CELL RECEPTOR BETA; CHAIN:	Ę;	FAB FRAGMENT; CHAIN:	ENDOTHELIAL GROWTH	FACTOR; CHAIN: V, W;	ANTIBODY R24 (LIGHT	CHAUN); CHAIN: A; ANTIRODY R24 (HEAVY	CHAIN); CHAIN: B;	IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B,
SeqFold Score							130.55								57.72			
PMF Score	0.31	0.17		0.88	060							0.92						0.93
Verify Score	-0.13	0.09		0.10	0.00	-						0.24						-0.12
PSI BLAST Score	5.4e-15	1.7e-20	\neg	5.1e-67	1e-66		3.4e-57			•	;	1.7e-68			1e-59			3.4e-69
End	168	213		180	180		681				1	081			189	-		180
Start AA	73	98		23	23	-	24					23			23			23
Chain ID	¥	A		-l	A		Ω				,	ے			۷			Ą
PDB ID	lfyv	Ifyx		1b2w	1b6d		1bd2					1bj1			1bz7			ldee
SEQ ID NO:	195	561		562	562		562				85	790			295			562

PDB annotation	RESOLUTION BINDING 2 OUTSIDE THE ANTIGEN COMBINING SITE SUPERANTIGEN FAB VH3 3 SPECIFICITY			COMPLEX (IMMUNORECEPTOR/IMMUNOGLOBU LIN) COMPLEX (IMMUNORECEPTOR/IMMUNOGLOBU LIN)	IMMUNOGLOBULIN IMMUNOGLOBULIN, AUTOANTIBODY, COLD AGGLUTININ, HUMAN IGM 2 FAB FRAGMENT	IMMUNE SYSTEM HUMAN TCR/PEPTIDE/MHC COMPLEX, HLA- A2, HTLV-1, TAX, TCR, T2 CELL RECEPTOR, IMMUNE SYSTEM	
	RESOLUTION THE ANTIGES SUPERANTIG			COMPLEX (IMMUNO) (IMMUNO) (IMMUNO)	IMMUNOGI IMMUNOGI AUTOANTII AGGLUTINI FRAGMENT	TCR/PE A2, HTI RECEP	VONOV
Coumpound	D, F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H;	IMMUNOGLOBULIN FAB FRAGMENT OF HUMANIZED ANTIBODY 4D5, VERSION 4 1FVD 3	IMMUNOGLOBULIN IMMUNOGLOBULIN FAB FRAGMENT (MC/PC\$603) IMCP 4	NIS ALPHA-BETA T-CELL RECEPTOR; CHAIN: A, B, C, D; H57 FAB; CHAIN: E, F, G, H	IGM KAPPA CHAIN V-III (KAU COLD AGGLUTININ); CHAIN: A, C; IGM FAB REGION IV- J(H4)-C (KAU COLD AGGLUTININ); CHAIN: B, D:	MHC CLASS I HLA-4; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE P6A; CHAIN: C; HMAN T-CELL RECEPTOR; CHAIN: D; HLA-A 0201; CHAIN: E:	A CONTOCT ONTA Y
SeqFold Score				115.80		126.38	
PMF Score		0.76	0.74		86.0		77.0
Verify Score		-0.05	-0.13		-0.18		000
PSI BLAST Score		6.8e-66	1.7e-66	5.1e-52	3.4e-67	3.4e-53	3 10-66
End AA		180	183	681	180	681	183
Start AA		23	23	24	27	24	23
Chain ID		¥	ı	٧	A	Ω	1
PDB ID		1fvd	Ітср	Infd	1qir	lqrn	Ishe
SEQ NO:		562	562	562	562	562	295

Chain ID A A I I I I I I I I I I I I I I I I	Start AA 24 24 25 25 25 83 83	AA AA 189 180 180 187 147	PSI BLAST Score 1.4c-57 1.7c-66 5.1c-68	Score 0.22 0.29 -0.09	Score 0.99 0.90 1.00	Score 118.15	Coumpound ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B; TRI.9 FAB; CHAIN: L, H; TRI.9 FAB; CHAIN: L, H; HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4 CATHEPSIN L: HEAVY CHAIN; CHAIN: A, C; CATHEPSIN L: LIGHT CATHEPSIN L: LIGHT CHAIN; CHAIN: B, D;	PDB annotation RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL IMMUNOGLOBULIN TR.1.9, ANTI- THYROID PEROXIDASE, AUTOANTIBODY, 2 IMMUNOGLOBULIN HYDROLASE II FRAGMENT, CD74 FRAGMENT CYSTEINE PROTEINASE, CATHEPSIN, MHC CLASS II, INVARIANT 2 CHAIN,
I A A	7 7	81	2.7e-26 1.1e-41	-0.31	1.00	136.01	INVARIANT CHAIN; CHAIN; I, J; CATHEPSIN L.: HEAVY CHAIN; CHAIN: A, C; CATHEPSIN L.: LIGHT CHAIN; CHAIN: B, D; INVARIANT CHAIN; CHAIN; CHAIN; CHAIN: I, J; HLA-DR ANTIGENS ASSOCIATED INVARIANT CHAIN; CHAIN: A, B, C; HLA-DR ANTIGENS	THYROGLOBULIN TYPE-1 DOMAIN HYDROLASE II FRAGMENT, CD74 FRAGMENT CYSTEINE PROTEINASE, CATHEPSIN, MHC CLASS II, INVARIANT 2 CHAIN, THYROGLOBULIN TYPE-1 DOMAIN MAJOR HISTOCOMPATIBILITY COMPLEX HLA CLASS II HISTOCOMPATIBILITY COMPLEX HLA CLASS II HISTOCOMPATIBILITY COMPLEX, ANTIGEN PROCESSING, 2 OLIGOMERIZATION, CHAPERONIN MAJOR HISTOCOMPATIBILITY

SEQ ID NO:	PDB ID	Chain ID	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									ASSOCIATED INVARIANT CHAIN; CHAIN: A, B, C;	COMPLEX HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, GAMMA MAJOR HISTOCOMPATIBILITY COMPLEX, ANTIGEN PROCESSING, 2
563	li: E	A	7	81	1.4e-24	-0.31	1.00		HLA-DR ANTIGENS ASSOCIATED INVARIANT CHAIN; CHAIN: A, B, C;	MAJOR HISTOCOMPATIBILITY COMPLEX HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, GAMMA MAJOR HISTOCOMPATIBILITY COMPLEX, ANTIGEN PROCESSING, 2 OLIGOMERIZATION, CHAPERONIN
998	lalh	A	427	512	1e-22	-0.45	0.11		QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN
268	laih	V	488	570	1.2e-21	0.10	-0.09		QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN
998	lalh	A	544	604	1.7e-17	0.30	-0.07		QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B,	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN
998	lard		544	574	0.0001	90.0	0.10		TRANSCRIPTION REGULATION YEAST	

PDB annotation		DNA-BINDING REGULATORY PROTEIN ATF-2; CRE BINDING PROTEIN, ATF-2, TRANSCRIPTIONAL ACTIVATION 2 DOMAIN, ZN FINGER	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	
Coumpound	TRANSCRIPTION FACTOR ADRI (RESIDUES 102 - 130) 1ARD 3 (AMINO. TERMINAL ZINC FINGER DOMAIN) (NMR, 10 STRUCTURES) 1ARD 4 (ADRIB) 1ARD 5	CRE-BPI; CHAIN: NULL;	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	TRANSCRIPTION REGULATION YEAST TRANSCRIPTION FACTOR ADRI (RESIDUES 130 - 159) IPAA 3 (PAPA - CARBOXY TERMINAL ZINC FINGER DOMAIN) MUTANT WITH IPAA 4 PRO 131
SeqFold Score						
PMF Score		99.0	-0.13	-0.20	0.11	0.51
Verify Score		-0.09	0.16	0.11	-0.21	-0.37
PSI BLAST Score		0.00034	6.8e-25	5.1e-11	1.2e-11	0.00085
End AA		570	604	339	570	572
Start AA		543	543	309	541	544
Chain ID			၁	Ŋ	Ö	
PDB ID		16hi	lmey	lmey	Ітеу	I paa
SEQ ID NO:		268	268	268	568	268

PDB annotation		ZINC FINGER TRANSCRIPTION FACTOR SP1; ZINC FINGER, TRANSCRIPTION ACTIVATION, SP1	COMPLEX (TRANSCRIPTION REGULATION/DNA) TFIIIA; 5S GENE; NMR, TFIIIA, PROTEIN, DNA, TRANSCRIPTION FACTOR, 5S RNA 2 GENE, DNA BINDING PROTEIN, ZINC FINGER, COMPLEX 3 (TRANSCRIPTION REGULATION/DNA)	ZINC FINGER DNA BINDING DOMAIN DNA BINDING MOTIF, ZINC FINGER DNA BINDING DOMAIN	TRANSCRIPTION REGULATION TRANSCRIPTION REGULATION, ADR1, ZINC FINGER, NMR	HEXAMERIZATION DOMAIN HEXAMERIZATION DOMAIN, ATPASE, TRANSPORT	CHAPERONE HSLV; HSLU CHAPERONE, HSLVU, CLPQY, AAA- ATPASE, ATP-DEPENDENT 2 PROTEOLYSIS, PROTEASOME	CELL CYCLE CDC6P; CDC6, CDC18, ORC1, AAA PROTEIN, DNA REPLICATION INITATION 2 FACTOR,
Coumpound	REPLACED BY ALA, PRO 133 REPLACED BY ALA, CYS 140 1PAA 5 REPLACED BY ALA (P131A,P133A,C140A) (NMR, 10 STRUCTURES) 1PAA 6	SPIF2; CHAIN: NULL;	TRANSCRIPTION FACTOR IIIA; CHAIN: A; SS RNA GENE; CHAIN: E, F;	SWIS; CHAIN: NULL;	ADRI; CHAIN: NULL;	N-ETHYLMALEIMIDE- SENSITIVE FUSION PROTEIN; CHAIN; A;	HEAT SHOCK PROTEIN HSLV; CHAIN: A, B, C, D; HEAT SHOCK PROTEIN HSLU; CHAIN: E, F;	CELL DIVISION CONTROL PROTEIN 6; CHAIN: A, B;
SeqFold Score								
PMF Score		0.80	-0.18	0.29	0.03	0.82	0.05	-0.05
Verify Score		0.07	0.18	0.19	0.26	0.29	-0.51	0.01
PSI BLAST Score		5.1e-10	6.8e-11	3.4e-06	8.5e-06	1.7e-13	3.46-12	1.9e-14
End		574	605	570	576	640	598	732
Start AA		544	544	543	544	489	477	490
Chain ID			¥.			Æ	ъ	Ą
PDB ID		1sp2	£11	pJzI	2adr	1d2n	1e94	1fnn
SEQ NO:		999	895	268	568	569	695	695

CLPY, LYSIS CLPY, CLPY, CLPY, CLPY,	, CLPY, LYSIS , CLPY, LYSIS S KINASE, ADP,	LYSIS LYSIS LYSIS LYSIS LYSIS SKINASE, ADP, LOOP LOOP LATASE, ACTIONS, Y	CLPY, LYSIS LYSIS CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLPY, CLOOP ADMAIN, AL AL	CLPY, LYSIS LYSIS CLPY,	CLPY, LYSIS LYSIS CLPY, LYSIS SKINASE, ADP, LOOP ADP, LOOP ADP, LOOP ADP, AL ROTEIN MAIN, AL ROTEIN	CLPY, LYSIS LYSIS LYSIS LYSIS SKINASE, ADP, LOOP LOOP LOOP LOOP LOOP AL RATASE, AL RAIN, AL ROTEIN MAIN, AL ROTEIN
CHAPERONE AAA-AIPASE, CLPY ATP-DEPENDENT PROTEOLYSIS CHAPERONE AAA-ATPASE, CLPY ATP-DEPENDENT PROTEOLYSIS TREANSEED ASE SHICKAATE PROTEOTATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE SHICKAATE TO ANSEED ASE TO ANSEED	CHAPERONE AAA-A IPASE, CLPY, ATP-DEPENDENT PROTEOL YSIS CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTEOL YSIS TRANSFERASE SHIKIMATE KINASE, PHOSPHORYL TRANSFER, ADP, SHIKIMATE 2 PATHWAY, P-LOOP PROTEIN, TRANSFERASE	CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTEOL YSIS CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTEOL YSIS TRANSFERASE SHIKIMATE KINASE, PHOSPHORYL TRANSFER, ADP, SHIKIMATE 2 PATHWAY, P-LOOP PROTEIN, TRANSFERASE HYDROLASE TETRATRICOPEPTIDE, TRP; HYDROLASE, PHOSPHATASE, PROTEIN-PROTEIN INTERACTIONS, TPR, 2 SUPER-HELIX, X-RAY	CHAPERONE AAA-ATPASE, CLP) ATP-DEPENDENT PROTEOLYSIS CHAPERONE AAA-ATPASE, CLP) ATP-DEPENDENT PROTEOLYSIS TRANSFERASE SHKIMATE KINAPHOSPHORYL TRANSFER, ADP, SHIKIMATE 2 PATHWAY, P-LOOF PROTEIN, TRANSFERASE HYDROLASE TETRATRICOPEPTI TRP; HYDROLASE, PHOSPHATAS PROTEIN-PROTEIN INTERACTIOI TPR, 2 SUPER-HELIX, X-RAY STRUCTURE CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPFAT, HSP90, 2 PROTEIN RIND	CHAPERONE AAA-A IPASE, CLPY, ATP-DEPENDENT PROTEOLYSIS CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTEOLYSIS TRANSFERASE SHIKIMATE KINASE, PHOSPHORYL TRANSFER, ADP, SHIKIMATE 2 PATHWAY, P-LOOP PROTEIN, TRANSFERASE HYDROLASE TETRATRICOPEPTIDE, TRP; HYDROLASE, PHOSPHATASE, PROTEIN-PROTEIN INTERACTIONS, TPR, 2 SUPER-HELIX, X-RAY STRUCTURE CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSP90, 2 PROTEIN BINDING CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSP90, 2 HSP70, PROTEIN REPEAT, HSC70, 2 HSP70, PROTEIN	CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTECLYSIS CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTECLYSIS TRANSFERASE SHIKIMATE KINAS PHOSPHORYL TRANSFER, ADP, SHIKIMATE 2 PATHWAY, P-LOOP PROTEIN, TRANSFERASE HYDROLASE, PHOSPHATASE PROTEIN-PROTEIN INTERACTIONS TRP, HYDROLASE, PHOSPHATASE CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSP90, 2 PROTEIN BINDIN CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSC70, 2 HSP70, PROTEIN BINDING CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSC70, 2 HSP70, PROTEIN BINDING CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSC70, 2 HSP70, PROTEIN	CHAPERONE AAA-A IPASE, CLPY, ATP-DEPENDENT PROTEOLYSIS CHAPERONE AAA-ATPASE, CLPY, ATP-DEPENDENT PROTEOLYSIS TRANSFERASE SHIKIMATE KINASE, PHOSPHORYL TRANSFER, ADP, SHIKIMATE 2 PATHWAY, P-LOOP PROTEIN, TRANSFERASE HYDROLASE TETRATRICOPEPTIDE, TRP; HYDROLASE, PHOSPHATASE, PROTEIN, TRANSFERASE CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSP90, 2 PROTEIN BINDING CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSC70, 2 HSP70, PROTEIN BINDING CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSC70, 2 HSP70, PROTEIN BINDING CHAPERONE HOP, TPR-DOMAIN, PEPTIDE-COMPLEX, HELICAL REPEAT, HSC70, 2 HSP70, PROTEIN BINDING SIGNALING PROTEIN PEROXISMORE SIGNALING PROTEIN PEROXISMORE RECEPTOR 1, PTS1-BP, PEROXIN-5, TETRATRICOPEPTIDE REPEAT, TPR, 7 HELICAL REPEAT
ATP-DEPENI	CHAPERONE ATP-DEPENI TRANSFERA PHOSPHORY SHIKIMATE PROTEIN, TH		ш	ш	tu l	ω
F VINIA CE.	SHIKIMATE KINASE; CHAIN: A, B;	SHIKIMATE KINASE; CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATASE 5; CHAIN: NULL;	SHIKIMATE KINASE; CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATASE 5; CHAIN: NULL; TPRZA-DOMAIN OF HOP; CHAIN: A; HSP90-PEPTIDE MEEVD: CHAIN: B:	SHIKIMATE KINASE; CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATASE 5; CHAIN: NULL; TPR2A-DOMAIN OF HOP; CHAIN: A; HSP90-PEPTIDE MEEVD; CHAIN: B; TPR1-DOMAIN OF HOP; CHAIN: A, B; HSC70- CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D;	SHIKIMATE KINASE; CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATASE 5; CHAIN: NULL; TPRZA-DOMAIN OF HOP; CHAIN: A; HSP90-PEPTIDE MEEVD; CHAIN: B; TPR1-DOMAIN OF HOP; CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D; TPR1-DOMAIN OF HOP; CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D; TPR1-DOMAIN OF HOP; CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D;	SHIKIMATE KINASE; CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATASE 5; CHAIN: NULL; TPR2A-DOMAIN OF HOP; CHAIN: A; HSP90-PEPTIDE MEEVD; CHAIN: B; TPR1-DOMAIN OF HOP; CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D; TPR1-DOMAIN OF HOP; CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D; PEROXISOMAL TARGETING SIGNAL 1 RECEPTOR; CHAIN: A, B; PTS1-CONTAINING PEPTIDE; CHAIN: A, B; PTS1-CONTAINING PEPTIDE; CHAIN: C, D;
TIVINITIE	CHAIN: A, B;	CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHAT 5; CHAIN: NULL;	CHAIN: A, B. SERINE/THR PROTEIN PH 5; CHAIN: NI TPR2A-DOM CHAIN: A; H MEEVD: CH	CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATA 5; CHAIN: NULL; TPR2A-DOMAIN OF HC CHAIN: A; HSP90-PEPT MEEVD; CHAIN: B; TPR1-DOMAIN OF HOP CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D;	CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATA 5; CHAIN: NULL; TPRZA-DOMAIN OF HC CHAIN: A; HSP90-PEPT MEEVD; CHAIN: B; TPR1-DOMAIN OF HOP CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D; TPR1-DOMAIN OF HOP CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D;	CHAIN: A, B; SERINE/THREONINE PROTEIN PHOSPHATA 5; CHAIN: NULL; 5; CHAIN: NULL; TPRZA-DOMAIN OF HO CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D TPRI-DOMAIN OF HOI CHAIN: A, B; HSC70- PEPTIDE; CHAIN: C, D PEROXISOMAL TARGETING SIGNAL TARGETING SIGNAL TARGETING SIGNAL TRECEPTOR; CHAIN: A, PTS1-CONTAINING PEPTIDE; CHAIN: A, PTS1-CONTAINING
1		0.48	0.10	0.10	0.10 0.00 0.72	0.10 0.00 0.72 0.45
3		0.38	0.28	0.38	0.38 0.28 -0.06	0.38 0.28 -0.06 0.07
		8.1e-05	8.1e-05	8.1e-05 0.00054 1.6e-05	8.1e-05 0.00054 1.6e-05 0.0027	8.1e-05 0.00054 1.6e-05 5.4e-05
}		170	170	170	170 192 233 233 143	170 192 193 233 143 150
<u>.</u>		20	50	50 110 1162	110 110	50 110 110 6
:			4	4 4	4 4 4	4 4 4 4
		1a17	lai7	la17 lelr lelw	lai7 lelr lelw lelw	lai7 lelw lelw lelw lfch
		270	570	570 570	570 570 570	570 570 570 570

SEQ ID NO:	PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
575	laox	А	991	363	3.4e-36	1.09	00.1		INTEGRIN ALPHA 2 BETA; CHAIN: A, B;	INTEGRIN INTEGRIN, CELL ADHESION, GLYCOPROTEIN
575	laox	V	991	366	3.4e-36			191.82	INTEGRIN ALPHA 2 BETA; CHAIN: A. B:	INTEGRIN INTEGRIN, CELL ADHESION, GLYCOPROTEIN
575	lauq		156	370	3.4e-40	0.21	0.99		AI DOMAIN OF VON	WILLEBRAND WILLEBRAND, BLOOD
								-	WILLEBRAND FACTOR; CHAIN: NIII.I.:	COAGULATION, PLATELET,
575	1ck4	A	169	361	1.9e-61	1.20	1.00		INTEGRIN ALPHA-1;	STRUCTURAL PROTEIN I-DOMAIN,
									CHAIN: A, B;	METAL BINDING, COLLAGEN, ADHESION
575	1ck4	¥	171	359	1.7e-35	1.12	1.00		INTEGRIN ALPHA-1;	STRUCTURAL PROTEIN I-DOMAIN,
									CHAIN: A, B;	METAL BINDING, COLLAGEN, ADHESION
575	Ifus	Ą	166	367	1e-37	0.47	1.00		IMMUNOGLOBULIN NMC-	IMMUNE SYSTEM VON WILLEBRAND
						•			4 IGG1; CHAIN: L;	FACTOR, GLYCOPROTEIN IBA
									IMMUNOGLOBULIN NMC-	(A:ALPHA) BINDING, 2 COMPLEX
								-	4 IGG1; CHAIN: H; VON	(WILLEBRAND/IMMUNOGLOBULIN),
									WILLEBRAND FACTOR;	BLOOD COAGULATION TYPE 3 2B
575	1005	4	391	350	5 10.15	1 14	6	 	AI BUA I BETA I WITEGBRE	CELL A DITECTON INTEGRAL OFFI
3	<u> </u>	€	801	900	3.46-43	01:10	8.		ALPHAI BETAT INTEGRIN; CHAIN: A; ALPHAI BETAT INTEGRIN: CHAIN: B:	CELL ADHESION INTEGRIN, CELL ADHESION
575	19c5	A	171	359	6.8e-36	1.16	1.00		ALPHAI BETAI INTEGRIN;	CELL ADHESION INTEGRIN, CELL
									CHAIN: A; ALPHAI BETAI INTEGRIN; CHAIN: B;	ADHESION
283	lalh	∢	201	283	5.4e-37	0.27	1.00		QGSR ZINC FINGER PEPTIDE; CHAIN: A;	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC
	•								DUPLEX	FINGER, DNA-BINDING PROTEIN
									OLIGONUCLEOTIDE	
									BINDING SITE; CHAIN: B, C:	
583	lalh	A	201	283	5.4e-37			68.06	QGSR ZINC FINGER	COMPLEX (ZINC FINGER/DNA)

	T							_	——	_										_
PDB annotation	COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN		COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING FROIEIN	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING PROTEIN		CONTRACTILE LIM DOMAIN, CRP,	NMR, MUSCLE DIFFERENTIATION, CONTRACTILE	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	CRYSTAL STRUCTURE. COMPLEX	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	Chystal stricture Colonies	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX
Coumpound	PEPTIDE; CHAIN: A; DUPLEX	OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	QGSR ZINC FINGER PEPTIDE; CHAIN: A;	OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	QGSR ZINC FINGER PEPTIDE; CHAIN: A;	DUPLEX	OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	CRP1; CHAIN: A;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	rnotelin; chain; C, f, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	PROTEIN; CHAIN: C, F, G;	
SeqFold Score								55.06						108.46						
PMF Score			0.93		0.88					1.00								1.00		
Verify Score			-0.05		0.10					0.70					-		_	0.58		
PSI BLAST Score			2.7e-24		5.1e-23			5.4e-13		3.4e-51	_			3.4e-51				6.8e-51		
End AA			296		309			315		161				198				225		
Start AA			229		229			112		911				116				144		
Chain ID			۷		A			Ą		O				ပ				၁		
PDB ID			lalh		lalh			1b8t		lmey				lmey				lmey		
SEQ ID NO:			583		583			283		583				583				583		

DNA; CHAIN: A, B, D, E; COMPLEX (ZINC FINGER/DNA) CONSENSUS ZINC FINGER, PROTEIN-DNA PROTEIN; CHAIN: C, F, G; INTERACTION, PROTEIN DESIGN, 2	++-		(ZIN	0.73 1.00 DNA: CHAIN: A. B. D. E.	N(Z)				
 	+			0.73 1.00 DNA: CHAIN: A. B. D. E.					
	-	DNA; CHAIN: A, B, D, E;	1.00 DNA; CHAIN: A, B, D, E;		1.00 DNA; CHAIN: A, B, D, E;	0.73 1.00 DNA; CHAIN: A, B, D, E;	1.4e-50 0.73 1.00 DNA; CHAIN: A, B, D, E;	253 1.4e-50 0.73 1.00 DNA; CHAIN: A, B, D, E;	172 253 1.4e-50 0.73 1.00 DNA; CHAIN: A, B, D, E;
Aug -									
CKYSIAL SIRUCIUKE, CUMPLEX (ZINC FINGER/DNA)	CKY	CKY							
┢	┢	DNA; CHAIN: A, B, D, E;	1.00 DNA; CHAIN: A, B, D, E;	0.37 1.00 DNA; CHAIN: A, B, D, E;	1.00 DNA; CHAIN: A, B, D, E;	0.37 1.00 DNA; CHAIN: A, B, D, E;	274 3.4e-46 0.37 1.00 DNA; CHAIN: A, B, D, E;	274 3.4e-46 0.37 1.00 DNA; CHAIN: A, B, D, E;	200 274 3.4e-46 0.37 1.00 DNA; CHAIN: A, B, D, E;
<u>~</u>									
FROIEIN; CHAIN: C, F, G; INTERACTION, PROTEIN DESIGN, 2 CRYSTAL. STRUCTURE COMPLEX									
(ZINC FINGER/DNA)	(ZIN	(ZIN	(ZIN	(ZIN					
		DNA; CHAIN: A, B, D, E;	0.65 DNA; CHAIN: A, B, D, E;	0.65 DNA; CHAIN: A, B, D, E;	-0.01 0.65 DNA; CHAIN: A, B, D, E;	1.4e-40 -0.01 0.65 DNA; CHAIN: A, B, D, E;	309 1.4e-40 -0.01 0.65 DNA; CHAIN: A, B, D, E;	228 309 1.4e-40 -0.01 0.65 DNA; CHAIN: A, B, D, E;	C 228 309 1.4e-40 -0.01 0.65 DNA; CHAIN: A, B, D, E;
PROTEIN: CHAIN: C. F. G. INTERACTION PROTEIN DESIGN 2									
(ZINC FINGER/DNA	(ZINC FI	(ZINC FI	(ZINC FI	(ZINC FI	(ZINC FI	(ZINC FI			
DNA; CHAIN: A, B, D, E; COMPLEX (ZINC FINGER/DNA) ZINC	-	DNA; CHAIN: A, B, D, E;	0.41 DNA; CHAIN: A, B, D, E;	DNA; CHAIN: A, B, D, E;	0.41 DNA; CHAIN: A, B, D, E;	1e-25 -0.33 0.41 DNA; CHAIN: A, B, D, E;	317 1e-25 -0.33 0.41 DNA; CHAIN: A, B, D, E;	256 317 1e-25 -0.33 0.41 DNA; CHAIN: A, B, D, E;	256 317 1e-25 -0.33 0.41 DNA; CHAIN: A, B, D, E;
~									
PROTEIN; CHAIN: C, F, G; INTERACTION, PROTEIN DESIGN, 2									
CRYSTAL STRUCTU (ZINC FINGER/DNA)	CRYSIA (ZINC F	CRYSIA (ZINC F	CRYSIA (ZINC F)	CRYSIA (ZINC F	CRYSIA (ZINC F	CRYSIA (ZINC F)	CRYSIA (ZINC F)	CRYSIA (ZINC F)	CRYSIA (ZINC F
		DNA: CUANIA B P.E.	000 PMA, CHANK A D P. F.	000 PMA, CHANK A D P. F.	031 000 DMA: CHAINLA B P. F.	1.2p. 30 0.31 0.00 DMA: CHAIN! A D F.	113 126.30 031 000 DMA: CHAINLA B D. E.	44 113 126-39 031 000 DNA CHAIN, A B D. E.	C 44 113 126-30 031 000 PMA CHANK A D P. F.
DNA; CHAIN: A, B, D, E; COMPLEX (ZINC FINGER/DNA) ZINC		DNA; CHAIN: A, B, D, E;	0.99 DNA; CHAIN: A, B, D, E;	DNA; CHAIN: A, B, D, E;	0.99 DNA; CHAIN: A, B, D, E;	0.31 0.99 DNA; CHAIN: A, B, D, E;	1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E;	113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E;	44 113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E;
		DNA; CHAIN: A, B, D, E;	0.99 DNA; CHAIN: A, B, D, E;	0.99 DNA; CHAIN: A, B, D, E;	0.31 0.99 DNA; CHAIN: A, B, D, E;	1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E;	113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E;	44 113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	C 44 113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER
\vdash	\vdash	DNA; CHAIN: A, B, D, E;	DNA; CHAIN: A, B, D, E;	DNA; CHAIN: A, B, D, E;	0.31 0.99 DNA; CHAIN: A, B, D, E;	1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E;	113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E; CONSENSIS ZING EINIGED	44 113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER	C 44 113 1.2e-39 0.31 0.99 DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER
DNA; CHAIN: A, B, D, E;	DNA; CHAIN: A, B, D, E;		66.0	66.0	0.31 0.99	1.26-39 0.31 0.99	11.3 1.26-39 0.31 0.99	44 113 1.2e-39 0.31 0.99	C 44 113 1.26-39 0.31 0.99
DNA; CHAIN: A, B, D, E;	DNA; CHAIN: A, B, D, E;		66.0	0.99	0.31 0.99	1.2e-39 0.31 0.99	113 1.2e-39 0.31 0.99	44 113 1.2e-39 0.31 0.99	C 44 113 1.2e-39 0.31 0.99
DNA; CHAIN: A, B, D	DNA; CHAIN: A, B, D		0.99	0.99	0.31 0.99	1.2e-39 0.31 0.99	113 1.2e-39 0.31 0.99	44 113 1.2e-39 0.31 0.99	C 44 113 1.2e-39 0.31 0.99
CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G; DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G; DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G; DNA; CHAIN: A, B, D, E;	CONSENSUS ZA PROTEIN; CHAI DNA; CHAIN: A, CONSENSUS ZI PROTEIN; CHAI CONSENSUS ZI PROTEIN; CHAI PROTEIN; CHAI		3 0.41	3 0.41	-0.01 0.65 -0.33 0.41	1.4e-40 -0.01 0.65 1e-25 -0.33 0.41 1.2e-39 0.31 0.99	8 309 1.4e-40 -0.01 0.65 5 317 1e-25 -0.33 0.41 113 1.2e-39 0.31 0.99	228 309 1.4e-40 -0.01 0.65 256 317 1e-25 -0.33 0.41 44 113 1.2e-39 0.31 0.99	C 228 309 1.4e-40 -0.01 0.65 C 256 317 1e-25 -0.33 0.41 C 44 113 1.2e-39 0.31 0.99
PROTEIN; DNA; CHA CONSENS; PROTEIN; DNA; CHA CONSENS! PROTEIN;	PROTEIN; DNA; CHA CONSENS PROTEIN; DNA; CHA CONSENSI PROTEIN; PROTEIN;		3 0.41	3 0.41	-0.01 0.65 -0.33 0.41 0.31 0.99	1.4e-40 -0.01 0.65 1e-25 -0.33 0.41 1.2e-39 0.31 0.99	309 1.4e-40 -0.01 0.65 317 1e-25 -0.33 0.41 113 1.2e-39 0.31 0.99	228 309 1.4e-40 -0.01 0.65 256 317 1e-25 -0.33 0.41 44 113 1.2e-39 0.31 0.99	C 228 309 1.4e-40 -0.01 0.65 C 256 317 1e-25 -0.33 0.41 C 44 113 1.2e-39 0.31 0.99
PROJ DNA, CONS, PROJ PROJ	PROJ DNA, CONS PROJ PROJ		3 0.41	3 0.41	-0.01 0.65	1.4e-40 -0.01 0.65 1e-25 -0.33 0.41 1.2e-39 0.31 0.99	309 1.4e-40 -0.01 0.65 317 1e-25 -0.33 0.41	228 309 1.4e-40 -0.01 0.65 256 317 1e-25 -0.33 0.41	C 228 309 1.4e-40 -0.01 0.65 C 256 317 1e-25 -0.33 0.41 C 44 113 1.2e-39 0.31 0.99
			3 0.41	3 0.41	-0.01 0.65 -0.33 0.41	1.4e-40 -0.01 0.65 1e-25 -0.33 0.41 1.2e-39 0.31 0.99	309 1.4e-40 -0.01 0.65 317 1e-25 -0.33 0.41 113 1.2e-39 0.31 0.99	228 309 1.4e-40 -0.01 0.65 256 317 1e-25 -0.33 0.41 44 113 1.2e-39 0.31 0.99	C 228 309 1.4e-40 -0.01 0.65 C 256 317 1e-25 -0.33 0.41 C 44 113 1.2e-39 0.31 0.99
		3.41		-0.37	-0.01	3.4e-46 0.37 1.4e-40 -0.01 1e-25 -0.33	309 1.4e-46 0.37 309 1.4e-40 -0.01 317 1e-25 -0.33	200 274 3.4e-46 0.37 228 309 1.4e-40 -0.01 256 317 1e-25 -0.33 44 113 1.2e-39 0.31	C 200 274 3.4e-46 0.37 C 228 309 1.4e-40 -0.01 C 256 317 1e-25 -0.33 C 44 113 1.2e-39 0.31

														_	_														
PDB annotation	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA INTERACTION PROTEIN DESIGN 2	CRYSTAL STRUCTURE, COMPLEX	(ZINC FINGER/DNA)	COMPLEX (TRANSCRIPTION PEGIT ATTONOMA) TEITA: 55 GENE:	NMR, TFIIIA, PROTEIN, DNA,	TRANSCRIPTION FACTOR, 5S RNA 2	GENE, DNA BINDING PROTEIN, ZINC	FINGER, COMPLEX 3	(IKANSCKIPTION KEGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION REGULATION/DNA) COMPLEX
Coumpound		DNA; CHAIN: A, B, D, E;	PROTEIN: CHAIN: C. F. G.			TRANSCRIPTION FACTOR	GENE; CHAIN: E, F;	•		•	OF W. I THINK I THE	IFIIIA; CHAIN: A, D; 5S RIBOSOMAL RNA GENE:	CHAIN: B, C, E, F;				TFIIIA; CHAIN: A, D; 5S	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;				TFIIIA; CHAIN: A, D; 5S	KIBOSOMAL KNA GENE;	CHAIN: B, C, E, F;				TFIIIA; CHAIN: A, D; 5S RIBOSOMAL RNA GENE:
SeqFold Score																							116.35						
PMF Score		1.00				0.11					000	78.0					1.00												1.00
Verify Score		0.27				-0.07					700	0.04					0.41												0.49
PSI BLAST Score		1.6e-10				1.2e-13					76 00 7	0.86-30					1.5e-34				-		le-38						1.2e-38
End AA		281				305					316	310					178						225						206
Start AA		254				229					146	£		,			43				-	1	09						61
Chain ID		Ð				Ψ					_	<					4						∢						Ą
PDB ID		Imey				Œ				_	1.6%	0111					1tf6		_			1	911						1tf6
SEQ ID NO:		583				283					502	S					583				_	1	283						583

			т			
PDB annotation	(TRANSCRIPTION REGULATION/DNA), RNA POLYMERASE III, 2 TRANSCRIPTION INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGII ATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX TRANSCRIPTION REGIT A TION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGIL ATTONDANA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGIT ATTON/DNA)	COMPLEX (TRANSCRIPTION REGULATIONDNA) YING-YANG 1;
Coumpound	CHAIN: B, C, E, F;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YY1; CHAIN: C; ADENO- ASSOCIATED VIRUS PS INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5
SeqFold Score					96.41	
PMF Score		1.00	1.00	1.00		1.00
Verify Score		0.60	0.38	0.35		0.19
PSI BLAST Score		5.4e-47	3.4e-35	2.7e-46	2.7e-46	5.4e-35
End AA		225	253	281	282	296
Start AA		114	152	170	174	198
Chain ID		၁	ပ	U	ပ	ပ
PDB CI		1ubd	lubd	lubd	lubd	lubd
SEQ NO:		583	583	583	583	583

1 # X	Start	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
	1		Score				INITIATOR ELEMENT DNA; CHAIN: A, B;	TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX TRANSCRIPTION 19 EGIT A TICKNINA A
308	١٣	309	1.7e-27	-0.21			YYI; CHAIN: C; ADENO-ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	COMPLEX (TRANSCRIPTION REGULATION DIA REGULATION DIA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGILI ATTONIONA)
141	12		8.5e-32	0.44	1.00		YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR, ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)
65 169	91		1.1e-47	0.37	1.00		YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS PS INITIATOR ELEMENT DNA; CHAIN: A, B;	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)
91	12	691	1.2e-34	0.48	1.00		YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS PS INITIATOR ELEMENT DNA; CHAIN: A, B;	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGUI ATION/DNA)
257 309	21	H	1.7e-08	-0.43	0.0		ADRI; CHAIN: NULL;	TRANSCRIPTION REGULATION

804 01	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
				Score					TRANSCRIPTION REGULATION,
✓		116	781	2 4e-58	0.40	1 00		ZINC EINGER DROTEIN	ADRI, ZINC FINGER, NMR
	-	?		2	?	3		GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
								CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA)
٧		124	252	5.1e-35	0.48	1.00		ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
								GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
								CHAIN: C, D;	LINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA)
∢		152	275	5.1e-32	0.55	66.0		ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
								GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
								CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)
⋖		180	308	6.8e-27	0.38	0.29		ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
								GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
								CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA-
_[1								BINDING PROTEIN/DNA)
⋖_		9	199	1.9e-62			104.54	ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
	-							GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
				_				CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA-
止		,		1					BINDING PROTEIN/DNA)
⋖		[9	661	1.9e-62	0.53	 0.		ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
								GLII; CHAIN: A; DNA;	FROI EIN/DNA) FIVE-FINGER GLI; GLI,
								CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA-
╧		87	107		\$ 6	5		י מחויים מי מיוטיומים ביומים	BINDING PROJEIN/DNA)
<		e e	8	0.86-34	0.42	3.		ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
								GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
		-			-			CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA-
Ŀ	T	00	200	+	30	00		· now our and ou of the	DINDLING FROIEIN/DINA)
<		00	177	70-34.7	0.04	1.00		ZINC FINGER PROTEIN	COMPLEX (DNA-BINDING
								GLII; CHAIN: A; DNA;	PROTEIN/DNA) FIVE-FINGER GLI; GLI,
					_			CHAIN: C, D;	ZINC FINGER, COMPLEX (DNA-
J			7						BINDING PROTEIN/DNA)

Chain Start ID AA	Star		End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
30 83		8		6.8e-23	-0.81	0.60		G PROTEIN GI ALPHA 1; CHAIN: A; G PROTEIN GI BETA 1; CHAIN: B; G PROTEIN GI GAMMA 2; CHAIN: G:	COMPLEX (GTP-BINDING/TRANSDUCER) SIGNAL TRANSDUCTION PROTEIN, GTPASE, WD40, RAS-LIKE, 2 COMPLEX (GTP-BINDING/TRANSDICER)
G 30 83		83		6.8e-23			57.14	G PROTEIN GI ALPHA 1; CHAIN: A; G PROTEIN GI BETA 1; CHAIN: B; G PROTEIN GI GAMMA 2; CHAIN: G;	COMPLEX (GTP- BINDING/TRANSDUCER) SIGNAL TRANSDUCTION PROTEIN, GTPASE, WD40, RAS-LIKE, 2 COMPLEX (GTP- BINDING/TRANSDUCER)
A 78 196		961		1.4e-18	0.49	0.35		GLYCINE N- METHYLTRANSFERASE; CHAIN: A. B. C. D:	TRANSFERASE METHYLTRANSFERASE
A 79 197		197		1.7e-07	-0.11	0.00		MJ0882; CHAIN: A;	STRUCTURAL GENOMICS HYPOTHETICAL PROTEIN, METHANOCOCCUS JANNASCHII
A 79 199		661		0.00017	0.09	0.16		FTSJ; CHAIN: A;	TRANSFERASE FTSJ METHYLTRANSFERASE; FTSJ, METHYLTRANSFERASE, ADOMET, ADENOSYL METHIONINE, HEAT 2 SHOCK PROTEINS, 23S RIBOSOMAL RNA
57 195		195		8.1e-09	0.28	0.04		CATECHOL O- METHYLTRANSFERASE; CHAIN: NULL;	TRANSFERASE (METHYLTRANSFERASE) COMT; TRANSFERASE, METHYLTRANSFERASE, NEUROTRANSMITTER DEGRADATION
78 196		196		1.4e-18	0.32	0.24		GLYCINE N- METHYLTRANSFERASE; CHAIN: A, B;	METHYLTRANSFERASE GNMT, S- ADENOSYL-L-METHIONINEN: GLYCINE METHYLTRANSFERASE

										_
PDB annotation		HORMONE RECEPTOR HORMONE RECEPTOR, INSULIN RECEPTOR FAMILY	MUSCLE PROTEIN CTNC; CARDIAC, MUSCLE PROTEIN, REGULATORY, CALCIUM BINDING	MUSCLE PROTEIN CTNC; CARDIAC, MUSCLE PROTEIN, REGULATORY, CALCIUM BINDING	CALCIUM-BINDING PROTEIN CALMODULIN CERIUM TR1C- DOMAIN, RESIDUES 1 - 75; CERUM- LOADED, CALCIUM-BINDING PROTEIN	HYDROLASE CALCINEURIN; HYDROLASE, PHOSPHATASE, IMMUNOSUPPRESSION	HYDROLASE CALCINEURIN; HYDROLASE, PHOSPHATASE, IMMUNOSUPPRESSION	MUSCLE CONTRACTION MUSCLE CONTRACTION, CALCIUM- ACTIVATED, TROPONIN, E-F HAND 2 CALCIUM-BINDING PROTEIN	CALCIUM-BINDING CALCIUM- BINDING, MYRISTOYLATION, NEURONAL SPECIFIC GUANYLATE 2 CYCLASE ACTIVATOR	CALCIUM-BINDING PROTEIN SNTNC;
Coumpound	VIRUS EQUINE HERPES VIRUS-1 (C3HC4, OR RING DOMAIN) 1CHC 3 (NMR, 1 STRUCTURE) 1CHC 4	INSULIN-LIKE GROWTH FACTOR RECEPTOR 1; CHAIN: A;	TROPONIN C; CHAIN: NULL;	TROPONIN C; CHAIN: NULL;	CALMODULIN; CHAIN: NULL;	SERINE/THREONINE PHOSPHATASE 2B; CHAIN: A, B;	SERINE/THREONINE PHOSPHATASE 2B; CHAIN: A, B;	TROPONIN C; CHAIN: A, B;	NEUROCALCIN DELTA; CHAIN: A, B;	N-TROPONIN C; CHAIN:
SeqFold Score				69.25			65.74		64.29	
PMF Score	0.01	0.04	0.92		0.65	0.83		66.0		69.0
Verify Score	-0.16	-0.22	0.58		0.41	0.58		0.67		0.50
PSI BLAST	5.4e-05	1.2e-05	6.8e-45	6.8e-45	1.7e-29	1.4e-40	1.4e-40	3.4e-25	3.4e-38	5.1e-26
End AA	179	169	168	170	68	175	179	16	185	93
Start AA	122	53	10		14	14	6	18	5	18
Chain ID		A				æ	В	A	ď	
PDB ID	1chc	ligr	laj4	laj4	lak8	1aui	laui	lavs	1bjf	1blq
SEQ	587	587	165	591	165	591	591	165	591	591

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PDB annotation	CALCIUM-BINDING, REGULATION, TROPONIN C, SKELETAL MUSCLE, 2 CONTRACTION																					CALCIUM-BINDING PROTEIN	CALMODULIN APO TRZC-DOMAIN;	, Hint
Coumpound	NULL;	CALCIUM-BINDING PROTEIN CALMODULIN COMPLEXED WITH	COMPLEASE WITH CALMODULIN-BINDING DOMAIN OF LCDM 3	CALMODULIN-	DEPENDENT PROTEIN KINASE II 1CDM 4	CALCIUM-BINDING PROTEIN CALMODITIN	COMPLEXED WITH	CALMODULIN-BINDING	DOMINIO OF ICDIM 3 CALMODULIN-	DEPENDENT PROTEIN	KINASE II 1CDM 4	CALCIUM-BINDING	PROTEIN CALMODULIN	CALCIUM-BINDING	PROTEIN CALMODULIN	(VEKTEBRATE) ICLL 3	CALCIUM-BINDING PROTEIN CALMODULIN	(VERTEBRATE) ICLL 3	CALCIUM-BINDING	PROTEIN CALMODULIN	(VERTEBRATE) ICLL 3	CALMODULIN	(VEKTEBKATE); ICMF 6 CHAIN: NULL: 1CMF 7	
SeqFold Score						65.81		_				-		75.45										
PMF Score		1.00										1.00				000	0.70		0.46		3	0.84		1
Verify Score		0.77										0.82				77.0	÷/.:		0.37			0 .07		T
PSI BLAST Score		3.4e-56				3.4e-56						8.5e-61		8.5e-61		1 20 26	07-27:1		1.7e-24		00 1	2.1e-28		
End		167				167						167		168		78	8		184		9	3	-	
Start AA		18				18						28		18		,	7		8		0	ŝ		
Chain ID		Ą				∢																		-
PDB ID		1cdm				1cdm				- 		- -]G]		=	=		Icil		3	IEE		
SEQ ID NO:		591				291						291		591		203	160		165		5	186		

SEQ	PDB	Chain	Start	End	. PSI	Verify	PMF	SeaFold	Coumnound	PDR annotation
e ë	e	<u>e</u>	ΑA	AA	BLAST Score	Score	Score	Score		
291	Idti	4	16	168	8.5e-43	0.71	1.00		CARDIAC TROPONIN C; CHAIN: A:	STRUCTURAL PROTEIN HELIX-TURN- HEI IX
591	lext	٧	16	168	5.1e-59	0.57	1.00		CALMODULIN; CHAIN: A;	METAL TRANSPORT CALMODULIN,
591	lexr	4	2	98	3.4e-25	0.51	1.00		CALMODULIN; CHAIN: A;	METAL TRANSPORT CALMODULIN,
591	lexr	Ą	88	184	3.4e-23	0.59	0.89		CALMODULIN; CHAIN: A;	METAL TRANSPORT CALMODULIN,
591	1731	A	95	169	3.4e-27	0.58	1.00		CALMODULIN; CHAIN: A;	TRANSPORT PROTEIN CALCIUM
										BINDING, EF HAND, FOUR-HELIX BUNDLE
591	16.5	∢	83	168	6.8e-20	-0.10	0.15		TROPONIN C; CHAIN: A;	CONTRACTILE PROTEIN TROPONIN
			-							C-IROPONIN I INTERACTION, CARDIAC, MUSCLE PROTEIN, 2
3			1							
186	2		-	186	3.4e-29	-		56.95	RECOVERIN; CHAIN:	CALCIUM-BINDING PROTEIN
									NULL;	CALCIUM-MYRISTOYL SWITCH,
591	Itcf		<u>s</u>	891	1.2e-47	0.75	8		TPODONINI C. CUANI.	CALCUIM-BINDING FROI BIN
			2	3		3	2		INCLUIN C; CHAIN:	CALCIUM-KEGULATED MUSCLE
										CONTRACTION, CALCIUM-BINDING.
										TROPONIN, E-F HAND, 2 OPEN
										CONFORMATION REGULATORY
										DOMAIN, CALCIUM-REGULATED 3
591	Itcf		2	98	5.1e-24	0.45	0.63		TROBONIN C. CHAIN:	MOSCLE CONTRACTION
							 !		NULL:	CALCIOM-NEOUGATED MUSCLE
		_								CONTRACTION CALCITIM-BINDING
										TROPONIN, E-F HAND, 2 OPEN
						-				CONFORMATION REGULATORY
					 ,					DOMAIN, CALCIUM-REGULATED 3
500	454	1	18		+	7				MUSCLE CONTRACTION
	1		3	184	8.3e-19	0.32	0.42		TROPONIN C; CHAIN: NULL;	CALCIUM-REGULATED MUSCLE CONTRACTION MUSCLE
									*	

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PDB annotation	CONTRACTION, CALCIUM-BINDING, TROPONIN, E-F HAND, 2 OPEN CONFORMATION REGULATORY DOMAIN, CALCIUM-REGULATED 3 MUSCLE CONTRACTION	CALCTUM-REGULATED MUSCLE CONTRACTION MUSCLE CONTRACTION, CALCTUM-BINDING, TROPONIN, E-F HAND, 2 OPEN CONFORMATION REGULATOR Y DOMAIN, CALCTUM-REGULATED 3 MUSCLE CONTRACTION	CALCIUM-BINDING PROTEIN EF- HAND I TNX 14	CALCIUM-BINDING PROTEIN EF- HAND 1TNX 14					
Coumpound		TROPONIN C; CHAIN: NULL;	TROPONIN C; 1TNX 4 CHAIN: NULL; 1TNX 5	TROPONIN C; ITNX 4 CHAIN: NULL; ITNX 5	CONTRACTILE SYSTEM PROTEIN TROPONIN C 1TOP 3	CONTRACTILE SYSTEM PROTEIN TROPONIN C 1TOP 3	CONTRACTILE SYSTEM PROTEIN TROPONIN C 1TOP 3	CALCIUM BINDING PROTEIN CALMODULIN (/TR=2=C\$ FRAGMENT COMPRISING RESIDUES 78 - 148 1TRC 3 OF THE INTACT MOLECULE) 1TRC	MUSCLE PROTEIN TROPONIN C (TRIC
SeqFold Score		77.47		69.47			78.36		
PMF Score			1.00		1.00	0.77		1.00	1.00
Verify Score			0.50		0.89	0.56		0.36	1.19
PSI BLAST Score		1.2e-47	5.1e-46	5.1e-46	6.8e-49	5.1c-24	6.8e-49	1.4e-27	3.4e-25
End AA		168	166	166	168	98	170	167	91
Start AA		٥.	18	6	18	2	9	83	18
Chain ID								V V	
PDB ID		ltcf	Itnx	1tnx	Itop	Itop	Itop	ltrc	ltr.
SEQ ID NO:		165	165	591	591	165	165	591	591

PDB annotation		CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALCIUM-BINDING PROTEIN CTNC; CARDIAC, MUSCLE, REGULATORY, CALCIUM-BINDING PROTEIN	MUSCLE PROTEIN CTNC; CARDIAC, MUSCLE PROTEIN, REGULATORY, CALCIUM BINDING	MUSCLE PROTEIN CTNC; CARDIAC, MUSCLE PROTEIN, REGULATORY, CALCIUM BINDING	CALCIUM-BINDING PROTEIN CALMODULIN CERIUM TRIC- DOMAIN, RESIDUES 1 - 75; CERIUM- LOADED, CALCIUM-BINDING PROTEIN	MUSCLE CONTRACTION MUSCLE
Coumpound	FRAGMENT) (APO FORM) (NMR, 1 STRUCTURE) 1TRF 3	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	TROPONIN C; CHAIN: NULL;	TROPONIN C; CHAIN: NULL;	TROPONIN C; CHAIN: NULL;	CALMODULIN; CHAIN: NULL;	TROPONIN C; CHAIN: A, B;
SeqFold Score			75.17					58.42		
PMF Score		1.00		0.99	0.63	0.59	0.98		0.65	0.99
Verify Score		0.75		0.15	0.29	0.22	0.85		0.41	0.67
PSI BLAST Score		1e-59	1e-59	1.4e-25	3.4e-23	1.2e-19	1e-40	1e-40	3.4e-30	1.4e-25
End		169	169	68	184	168	157	159	8	16
Start AA		15	16	2	87	16	12	1	14	18
Chain ID		A	A	A	Ą					A
PDB ID		lvrk	Ivrk	lvrk	lvrk	3ctn	laj4	laj4	lak8	lavs
SEQ NO:		591	591	591	591	591	591	291	165	591

														_							
PDB annotation	CONTRACTION, CALCIUM- ACTIVATED, TROPONIN, E-F HAND 2 CALCIUM-BINDING PROTEIN	CALCIUM-BINDING PROTEIN SNTNC; CALCIUM-BINDING, REGULATION, TROPONIN C, SKELETAL MUSCLE, 2	MUSCLE PROTEIN MDE; MUSCLE PROTEIN																		
Coumpound		N-TROPONIN C; CHAIN: NULL;	MYOSIN; CHAIN: A, B, C, D, E, F, G, H;	CALCIUM-BINDING	COMPLEXED WITH	CALMODULIN-BINDING DOMAIN OF ICDM 3	CALMODULIN-	DEPENDENT PROTEIN KINASE II 1CDM 4	CALCIUM-BINDING PROTEIN CALMODULIN	COMPLEXED WITH	DOMAIN OF ICDM 3	CALMODULIN-	DEPENDENT PROTEIN KINASE II 1CDM 4	CALCIUM-BINDING	PROTEIN CALMODULIN	CALCIUM-BINDING	PROTEIN CALMODULIN	(VERTEBRATE) 1CLL 3	CALCIUM-BINDING	PROTEIN CALMODULIN	(VEKIEBKAIE) ICLL 3
SeqFold Score									60.24				•			71.03			_		
PMF Score		69:0	0.99	1.00										1.00					0.90		
Verify Score		0:50	0.73	0.72										0.75					0.44		
PSI BLAST Score		5.1e-26	1.2e-33	5.1e-54	-				5.1e-54					le-57		1e-57			6.8e-26		
End		83	159	157	·				157					157		158			98		
Start AA		81	19	18					19					18		19			_		
Chain ID			В	Ą					4												
PDB ID		pld1	lbr1	1cdm					1cdm					Icli		le!			JCII		1
SEQ NO:		591	591	291					291					591		591			591		

<u>ු</u>	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
E S	8	A	AA	AA	BLAST Score	Score	Score	Score		
591	1dtl	٧	12	157	1.4e-40	69.0	1.00		CARDIAC TROPONIN C; CHAIN: A;	STRUCTURAL PROTEIN HELIX-TURN-HELIX
591	lexr	А	81	158	1.7e-55	05.0	1.00		CALMODULIN; CHAIN: A;	METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER
591	lexr	A	1	98	3.4e-23	0.53	0.53		CALMODULIN; CHAIN: A;	METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER
591	ltcf		12	158	1.7e-44			68.07	TROPONIN C; CHAIN:	CALCIUM-REGULATED MUSCLE
								-	NOLL;	CONTRACTION MUSCLE CONTRACTION, CALCIUM-BINDING,
	-									TROPONIN, E-F HAND, 2 OPEN
										CONFORMATION REGULATORY DOMAIN, CALCITM-REGULATED 3
591	Itcf		18	156	1.7e-44	0.28	1.00		TROPONIN C; CHAIN:	CALCIUM-REGULATED MUSCLE
									NULL;	CONTRACTION MUSCLE
										CONTRACTION, CALCIUM-BINDING,
										TROPONIN, E-F HAND, 2 OPEN
										DOMAIN, CALCIUM-REGULATED 3
7										MUSCLE CONTRACTION
291	Itnx		12	156	3.4e-43			63.93	TROPONIN C; 1TNX 4	CALCIUM-BINDING PROTEIN EF-
1									CHAIN: NULL; 1TNX 5	HAND 1TNX 14
291	Itmx		18	156	3.4e-43	0.51	0.94	,	TROPONIN C; 1TNX 4 CHAIN: NULL; 1TNX 5	CALCIUM-BINDING PROTEIN EF. HAND 1TNX 14
591	Itop		81	156	1.7e-45	89.0	1.00		CONTRACTILE SYSTEM	
									PROTEIN TROPONIN C	
7									1TOP 3	i
591	Itop		1	98	3.4e-23	0.48	0.25		CONTRACTILE SYSTEM	
									PROTEIN TROPONIN C	
7									1TOP 3	
 	 do		9	159	1.7e-45			69.75	CONTRACTILE SYSTEM PROTEIN TROPONIN C	
7									1TOP 3	
591	별		18	91	1.4e-25	1.19	1.00		MUSCLE PROTEIN	

SEQ ID NO:	PDB ID	Chain 1D	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									TROPONIN C (TRIC FRAGMENT) (APO FORM) (NMR, I STRUCTURE) ITRF 3	
591	1vrk	¥	15	157	8.5e-56	0.71	86.0		CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTINE)
591	lvrk	¥	17	159	8.5e-56			69.15	CALMODULN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)
591	lvrk	Ą	1	68	1.7e-24	0.30	0.70		CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)
591	2mys	В	20	145	1.7e-24	-0.16	0.34		MYOSIN; CHAIN: A, B, C;	MUSCLE PROTEIN MUSCLE PROTEIN, MYOSIN SUBFRAGMENT-1, MYOSIN HEAD, 2 MOTOR PROTEIN
592	laj4		10	168	6.8e-45	0.58	0.92		TROPONIN C; CHAIN:	MUSCLE PROTEIN CTNC; CARDIAC,
592	laj4			170	6.8e-45			69.25	TROPONIN C; CHAIN: NULL;	CALCIUM BINDING MUSCLE PROTEIN, REGULATORY, MUSCLE PROTEIN, REGULATORY,
592	lak8		14	68	1.7e-29	0.41	0.65		CALMODULIN; CHAIN: NULL;	CALCIUM-BINDING PROTEIN CALCHUM-BINDING PROTEIN CALMODULIN CERIUM TRIC- DOMAIN, RESIDUES 1 - 75; CERIUM- LOADED, CALCIUM-BINDING PROTEIN
592	laui	В	14	175	1.4e-40	0.58	0.83		SERINE/THREONINE PHOSPHATASE 2B; CHAIN: A, B;	HYDROLASE CALCINEURIN; HYDROLASE, PHOSPHATASE, IMMUNOSUPPRESSION

PDB annotation	HYDROLASE CALCINEURIN; HYDROLASE, PHOSPHATASE, IMMUNOSUPPRESSION	MUSCLE CONTRACTION MUSCLE CONTRACTION, CALCIUM- ACTIVATED, TROPONIN, E-F HAND 2 CAI CII M-BINDING PROTEIN	CALCIUM-BINDING CALCIUM- BINDING, MYRISTOYLATION, NEURONAL SPECIFIC GUANYLATE 2 CYCLASE ACTIVATOR	CALCIUM-BINDING PROTEIN SNTNC; CALCIUM-BINDING, REGULATION, TROPONIN C, SKELETAL MUSCLE, 2 CONTRACTION			
Coumpound	SERINE/THREONINE PHOSPHATASE 2B; CHAIN: A, B;	TROPONIN C; CHAIN: A, B;	NEUROCALCIN DELTA; CHAIN: A, B;	N-TROPONIN C; CHAIN: NULL;	CALCIUM-BNDING PROTEIN CALMODULIN COMPLEXED WITH CALMODULIN-BINDING DOMAIN OF 1CDM 3 CALMODULIN- DEPENDENT PROTEIN KINASE II 1CDM 4	CALCIUM-BINDING PROTEIN CALMODULIN COMPLEXED WITH CALMODULIN-BINDING DOMAIN OF ICDM 3 CALMODULIN- DEPENDENT PROTEIN KINASE II ICDM 4	CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) ICLL 3
SeqFold Score	65.74		64.29			65.81	
PMF Score		0.99		69.0	1.00		1.00
Verify Score		0.67		0.50	0.77		0.82
PSI BLAST Score	1.4e-40	3.4e-25	3.4e-38	5.1e-26	3.4e-56	3.46-56	8.5e-61
End	179	91	581	93	167	191	167
Start AA	6	18	٧,	18	18	18	18
Chain ID	В	A	V V		Y .	V	
PDB ID	laui	lavs	lbjf	1blq	lcdm	lcdm	Icll
SEQ ID NO:	592	292	592	592	592	592	592

Coumpound PDB annotation		NDING LMODULIN TE) ICLL 3	INDING LMODULIN TE) ICLL 3	JLIN L3		OPONIN C; STRUCTURAL PROTEIN HELIX-TURN-HELIX	CALMODULIN; CHAIN: A; METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER	CALMODULIN; CHAIN: A; METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER	CALMODULIN; CHAIN: A; METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER	CALMODULIN; CHAIN: A; TRANSPORT PROTEIN CALCIUM BINDING, EF HAND, FOUR-HELIX BUNDLE	CONTRACTILE PROTEIN TROPONIN C-TROPONIN I INTERACTION, CARDIAC, MUSCLE PROTEIN, 2 CALCIUM BINDING PROTEIN	CHAIN: CALCIUM-BINDING PROTEIN CALCIUM-MYRISTOYL SWITCH, CALCUIM-BINDING PROTEIN	TALL DEGIT ATEN MISCIE
		CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) 1CLL 3	CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) ICLL 3	CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) ICLL 3	CALMODULIN (VERTEBRATE); 1CMF 6 CHAIN: NULL; 1CMF 7	CARDIAC TROPONIN C; CHAIN: A;	CALMODULI	CALMODULI	CALMODULI	CALMODULI	TROPONIN C; CHAIN: A;	RECOVERIN; CHAIN: NULL;	TROPONIN C; CHAIN:
SeqFold	Score	75.45										56.95	
PMF	Score		0.98	0.46	0.84	1.00	1.00	1.00	0.89	1.00	0.15		1.00
Verify	Score		0.74	0.37	0.07	0.71	0.57	0.51	0.59	0.58	-0.10		0.75
PSI	BLAST Score	8.5e-61	1.2e-26	1.7e-24	5.1e-28	8.5e-43	5.1e-59	3.4e-25	3.4e-23	3.4e-27	6.8e-20	3.4e-29	1.2e-47
End	¥	168	98	184	169	168	168	98	184	169	168	186	168
Start	ΑĄ	18		06	68	16	16	2	88	95	83	_	18
Chain	<u>e</u>					4	∀	A	4	4	A		
PDB	<u>e</u>	101 1101	1cII	Icli	1cmf	1 g E	lexr	lexr	lexr	1771	165	1iku	1tcf
SEQ	АŞ	592	592	592	592	592	592	292	292	592	592	592	592

	7															
PDB annotation	CONFORMATION REGULATORY DOMAIN, CALCIUM-REGULATED 3 MUSCLE CONTRACTION	CALCIUM-REGULATED MUSCLE CONTRACTION MUSCLE CONTRACTION, CALCIUM-BINDING, TROPONIN, E-F HAND, 2 OPEN	CONFURMATION REGULATORY DOMAIN, CALCIUM-REGULATED 3 MUSCLE CONTRACTION	CALCIUM-REGULATED MUSCLE CONTRACTION MUSCLE CONTRACTION CALCILM-BINDING	TROPONIN, E-F HAND, 2 OPEN CONFORMATION REGULATORY	DOMAIN, CALCIUM-REGULATED 3 MUSCLE CONTRACTION	CALCIUM-REGULATED MUSCLE CONTRACTION MUSCLE	CONTRACTION, CALCIUM-BINDING,	CONFORMATION REGULATORY	DOMAIN, CALCIUM-REGULATED 3 MUSCLE CONTRACTION	CALCIUM-BINDING PROTEIN EF. HAND 1TNX 14	CALCIUM-BINDING PROTEIN EF-				
Coumpound		TROPONIN C; CHAIN: NULL;		TROPONIN C; CHAIN: NULL;			TROPONIN C; CHAIN: NULL;				TROPONIN C; 1TNX 4 CHAIN: NULL: 1TNX 5	TROPONIN C; 1TNX 4 CHAIN: NIII.1: 1TNX 5	CONTRACTILE SYSTEM	PROTEIN TROPONIN C	CONTRACTILE SYSTEM	PROTEIN TROPONIN C 1TOP 3
SeqFold Score							77.47					69.47		•		
PMF Score				0.42							1.00		1.00		0.77	
Verify Score		0.45		0.32							0.50		68.0		0.56	
PSI BLAST Score		5.1e-24		8.5e-19			1.2e-47				5.1e-46	5.1e-46	6.8e-49		5.1e-24	
End		98		184			168		,		166	166	168		98	
Start		7		8			6				8	6	18		2	
Chain 1D								- 								
PDB ID		ltcf		Itef			Itcf				Itnx	Itnx	Itop		Itop	
SEQ ID NO:		292		292			292				592	592	592		292	

PDB annotation				CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING	CALCIUM-BINDING PROTEIN CTNC; CARDIAC, MUSCLE, REGULATORY, CALCIUM-BINDING PROTEIN
Coumpound	CONTRACTILE SYSTEM PROTEIN TROPONIN C ITOP 3	CALCIUM BINDING PROTEIN CALMODULIN (/TR=2=C\$ FRAGMENT COMPRISING RESIDUES 78 - 148 1TRC 3 OF THE INTACT MOLECULE) 1TRC 4	MUSCLE PROTEIN TROPONIN C (TRIC FRAGMENT) (APO FORM) (NMR, I STRUCTURE) ITRF 3	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	TROPONIN C; CHAIN: NULL;
SeqFold Score	78.36		·		75.17			
PMF Score		1.00	1.00	1.00		66.0	0.63	0.59
Verify Score		0.36	1.19	0.75		0.15	0.29	0.22
PSI BLAST	6.8e-49	1.4e-27	3.4e-25	1e-59	1e-59	1.4e-25	3.4e-23	1.2e-19
End	170	167	16	169	169	68	184	168
Start AA	9	93	18	15	91	2	<i>L</i> 8	91
Chain ID		∢		A	Ą	¥	Y .	
ag a	Itop	1trc	弄	lvrk	lvrk	lvrk	Ivrk	3ctn
SEQ D	592	592	592	592	292	592	592	592

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PDB annotation		MUSCLE PROTEIN CTNC; CARDIAC, MUSCLE PROTEIN, REGULATORY, CALCIUM BINDING	MUSCLE PROTEIN CTNC; CARDIAC, MUSCLE PROTEIN, REGULATORY, CALCIUM BINDING	CALCIUM-BINDING PROTEIN	CALMODULIN CERIUM TRIC- DOMAIN, RESIDUES 1 - 75; CERIUM- LOADED, CALCIUM-BINDING	MUSCLE CONTRACTION MUSCLE	CONTRACTION, CALCIOM: ACTIVATED, TROPONIN, E-F HAND 2 CALCIUM-BINDING PROTEIN	CALCIUM-BINDING PROTEIN SNTNC;	CALCIUM-BINDING, REGULATION,	IROPONIN C, SKELETAL MUSCLE, 2 CONTRACTION	MUSCLE PROTEIN MDE; MUSCLE PROTEIN										
Coumpound		TROPONIN C; CHAIN: NULL;	TROPONIN C; CHAIN: NULL;	CALMODULIN; CHAIN:	NULL;	TROPONIN C; CHAIN: A, B;		N-TROPONIN C; CHAIN:	NULL;		MYOSIN; CHAIN: A, B, C, D, E, F, G, H:	CALCIUM-BINDING	PROTEIN CALMODULIN COMPLEXED WITH	CALMODULIN-BINDING	CALMODULIN-	DEPENDENT PROTEIN	KINASE II 1CDM 4	CALCIUM-BINDING	COMPLEXED WITH	CALMODULIN-BINDING	DOMAIN OF ICDM 3
SeqFold	Score		58.42															60.24			
PMF	SCOLE	0.98		0.65		0.99		69.0			0.99	1.00									
Verify	SCOIC	0.85		0.41		0.67		0.50			0.73	0.72									
PSI RIAST	Score	1e-40	Ie-40	3.4e-30		1.4e-25		5.1e-26			1.2e-33	5.1e-54						5.1e-54			
End		157	159	68		16		93			159	157						157			
Start		12		14		18		18			61	18						6			
Chain TD	9					Y					В	A						⋖			
PDB ID		1aj4 -	1aj4	1ak8		lavs		1blq			lbrl	Icdm						E G M			
SEQ	NO.	592	592	292		592		592			592	592					18	760			

PDB annotation					STRUCTURAL PROTEIN HELIX-TURN-HELIX	METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER	METAL TRANSPORT CALMODULIN, HIGH RESOLUTION, DISORDER	CALCIUM-REGULATED MUSCLE CONTRACTION MUSCLE CONTRACTION, CALCIUM-BINDING, TROPONIN, E-F HAND, 2 OPEN CONFORMATION REGULATORY DOMAIN CALCIUM-BEGILATED 3	MUSCLE CONTRACTION CALCIUM-REGULATED MUSCLE CONTRACTION MUSCLE	CONTRACTION, CALCIOM-BINDING, TROPONIN, E-F HAND, 2 OPEN CONFORMATION REGULATORY DOMAIN, CALCIUM-REGULATED 3 MISCLE CONTRACTION	CALCIUM-BINDING PROTEIN EF. HAND 1TNX 14
Coumpound	CALMODULIN- DEPENDENT PROTEIN KINASE II 1CDM 4	CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) 1CLL 3	CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) ICLL 3	CALCIUM-BINDING PROTEIN CALMODULIN (VERTEBRATE) ICLL 3	CARDIAC TROPONIN C; CHAIN: A;	CALMODULIN; CHAIN: A;	CALMODULIN; CHAIN: A;	TROPONIN C; CHAIN: NULL;	TROPONIN C; CHAIN: NULL;		TROPONIN C; 1TNX 4 CHAIN: NULL; 1TNX 5
SeqFold Score			71.03					68.07			63.93
PMF Score		1.00		0.90	1.00	1.00	0.53		1.00		
Verify Score		0.75		0.44	69.0	0.50	0.53		0.28		
PSI BLAST Score		le-57	le-57	6.8e-26	1.4e-40	1.7e-55	3.4e-23	1.7e-44	1.7e-44		3.4e-43
End AA		157	158	98	157	158	98	158	156		156
Start		18	19	-	12	18	-	12	18		12
Chain ID					٧	Ą	Ą				
PDB CI		Icli	ıcli	Icil	1dtl	lexr	lexr	ltef	ltcf		ltnx
SEQ EQ		292	592	592	592	292	292	592	592		592

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
U) NO:	ar	a	AA	AA	BLAST Score	Score	Score	Score		
592	Itnx		18	156	3.4e-43	0.51	0.94		TROPONIN C; 1TNX 4 CHAIN: NULL; 1TNX 5	CALCIUM-BINDING PROTEIN EF- HAND ITNX 14
592	ltop		18	156	1.7e-45	89.0	1.00		CONTRACTILE SYSTEM PROTEIN TROPONIN C 1TOP 3	
592	ltop		1	98	3.4e-23	0.48	0.25		CONTRACTILE SYSTEM PROTEIN TROPONIN C 1TOP 3	
592	Itop		9	159	1.7e-45			69.75	CONTRACTILE SYSTEM PROTEIN TROPONIN C 1TOP 3	
592	ltrf		18	91	1.4e-25	1.19	1.00		MUSCLE PROTEIN TROPONIN C (TRIC FRAGMENT) (APO FORM) (NMR, I STRUCTURE) ITRF 3	
592	lvrk	A	51	157	8.5e-56	0.71	0.98		CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)
592	lvrk	V V	17	159	8.5e-56			69.15	CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)
592	lvrk	¥	1	68	1.7e-24	0.30	0.70		CALMODULIN; CHAIN: A; RS20; CHAIN: B;	CALMODULIN, CALCIUM BINDING, HELIX-LOOP-HELIX, SIGNALLING, 2 COMPLEX(CALCIUM-BINDING PROTEIN/PEPTIDE)
592	2mys	В	50	145	1.7e-24	-0.16	0.34		MYOSIN; CHAIN: A, B, C;	MUSCLE PROTEIN MUSCLE PROTEIN, MYOSIN SUBFRAGMENT-1, MYOSIN HEAD, 2 MOTOR PROTEIN
594	1qsa	4	24	114	0.0019	-0.04	0.01		SOLUBLE LYTIC TRANSGLYCOSYLASE	TRANSFERASE ALPHA-SUPERHELIX, TRANSFERASE

PDB annotation		ISOMERASE ISOMERASE, MUTASE, INTRAMOLECULAR TRANSFERASE	PLANT PROTEIN TWO HOMOLOGOUS	GLYCOPROTEIN GLYCOPROTEIN	REPLICATION DNA DOUBLE-STRAND BREAK REPAIR, ABC-ATPASE	ACTIN-BINDING PROTEIN ACTIN- BINDING PROTEIN, CALCIUM- BINDING PHOSPHORYI ATON	STRICT ID AT BROTTENIOAT BONIEN	HOMOLOGY, ACTIN BINDING,	SINUCIONAL PROTEIN	STRUCTURAL PROTEIN CALPONIN HOMOLOGY, ACTIN BINDING, STRUCTURAL PROTEIN	ACTIN-BINDING CALPONIN	HOMOLOGY (CH) DOMAIN;	FILAMENTOUS ACTIN-BINDING	DOMAIN, CYTOSKELETON	ACTIN-BINDING CALPONIN	HOMOLOGY (CH) DOMAIN;	FILAMENTOUS ACTIN-BINDING	TRANSMEN OF TOSAELEION	COLICIN. BACTERIOCIN. ION	CHANNEL FORMATION,	I KAINSIMEIMIBIKAINE 2 PROTEIIN
Coumpound	SLT70; CHAIN: A;	METHYLMALONYL-COA MUTASE; CHAIN: A, B, C, D;	AGGLUTININ ISOLECTIN	LAMININ; CHAIN; NULL;	RAD50 ABC-ATPASE; CHAIN: A, C; RAD50 ABC- ATPASE; CHAIN: B, D;	T-FIMBRIN; CHAIN: NULL;	I TTP OPERIOR CHAIN! A B.	(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)		UTROPHIN; CHAIN: A, B;	SPECTRIN BETA CHAIN;	CHAIN: A;			SPECTRIN BETA CHAIN;	CHAIN: A;		COLICINITA: CHARA.	COLICIN IA; CHAIN: NULL:		
SeqFold Score	•									74.00	81.12					-					
PMF Score		0.31	0.07	0.21	0.46	0.47	9	3				•			1.00			000	07:0-		
Verify Score		0.24	0.98	0.91	-0.08	0.72	0.70	```							0.95			017	 		
PSI BLAST Score		2.7e-06	0.00014	0.00016	1e-26	5.1e-26	10.35			le-35	1.5e-43				1.5e-43			5 10 11	11-21-1		
End AA		114	83	83	186	481	480	}		483	486				486			260	3		7
Start AA		24	31	=	51	371	375			377	378				379		*******	53	3		
Chain ID		Ą	Ą		V		A			∢	A				¥	-					1
PDB ID		Ireq	lehd	1Klo	112m	laoa	1hhd			lohd	1bkr				Ibkr			::-			
SEQ B Si Si		594	595	595	009	604	809			604	604				604	_		604	3		

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation	
	a	a	AA	AA	BLAST Score	Score	Score	Score			
604	ldxx	K	373	490	1.7e-37	0.58	1.00		DYSTROPHIN; CHAIN: A, B, C, D;	STRUCTURAL PROTEIN DYSTROPHIN, MUSCULAR DYSTROPHY, CALPONIN HOMOLOGY DOMAIN, 2 ACTIN-BINDING, UTROPHIN	
604	lez3	A	109	239	1.6e-08	0.20	-0.17		SYNTAXIN-1A; CHAIN: A, B, C;	ENDOCYTOSIS/EXOCYTOSIS SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX BUNDLE	
604	lez3	A	77	192	5.4e-09	0.12	-0.19		SYNTAXIN-1A; CHAIN: A, B, C;	ENDOCYTOSIS/EXOCYTOSIS SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX BUNDLE	
604	lez3	¥	16	212	1.1e-08	0.31	-0.19		SYNTAXIN-1A; CHAIN: A, B, C;	ENDOCYTOSIS/EXOCYTOSIS SYNAPTOTAGMIN ASSOCIATED 35 KDA PROTEIN, P35A, THREE HELIX BUNDLE	
604	lqag	A	373	485	1.2e-36	0.70	1.00		UTROPHIN ACTIN BINDING REGION; CHAIN: A, B;	STRUCTURAL PROTEIN CALPONIN HOMOLOGY DOMAIN, DOMAIN SWAPPING, ACTIN BINDING, 2 UTROPHIN, DYSTROPHIN, STRUCTURAL PROTEIN	
604	1qu2	A	88	260	1.1e-09	0.11	-0.20		ISOLEUCYL-TRNA SYNTHETASE; CHAIN: A; ISOLEUCYL-TRNA; CHAIN: T;	LIGASE/RNA ISOLEUCINETRNA LIGASE, ILERS; PROTEIN-RNA COMPLEX, METAL IONS, EDITING TRNA SYNTHETASE, 2 DOUBLE- SIEVE	
604	lquu	A	63	259	5.4e-15	0.11	-0.19		HUMAN SKELETAL MUSCLE ALPHA-ACTININ 2; CHAIN: A;	CONTRACTILE PROTEIN TRIPLE- HELIX COILED COIL, CONTRACTILE PROTEIN	_
604	lreq.	A	_	261	2.7e-21	0.17	-0.18		METHYLMALONYL-COA MUTASE; CHAIN: A, B, C, D;	ISOMERASE ISOMERASE, MUTASE, INTRAMOLECULAR TRANSFERASE	
604	2trc	Ы	167	262	1.9e-09	0.18	-0.19		TRANSDUCIN; CHAIN: B,	COMPLEX	_

PDB annotation	P; (TRANSDUCER/TRANSDUCTION) GT BETA-GAMMA; MEKA, PP33; PHOSDUCIN, TRANSDUCIN, BETA- GAMMA, SIGNAL TRANSDUCTION, 2 REGULATION, PHOSPHORYLATION, G PROTEINS, THIOREDOXIN, 3 VISION, MEKA, COMPLEX (TRANSDUCER/TRANSDUCTION)		PHOSPHOLIPID DEGRADATION, VIRULENCE FACTOR OF 2 HUMAN PATHOGEN	HYDROLASE PI-PLC; HYDROLASE, PHOSPHOLIPID DEGRADATION, VIRULENCE FACTOR OF 2 HUMAN PATHOGEN	 	
Coumpound	G; PHOSDUCIN; CHAIN: P;	TRANSDUCIN; CHAIN: B, G; PHOSDUCIN; CHAIN: P;	PHOSPHATIDYLINOSITOL -SPECIFIC PHOSPHOLIPASE C; CHAIN: NULL;	PHOSPHATIDYLINOSITOL -SPECIFIC PHOSPHOLIPASE C; CHAIN: NULL:	PHOSPHATIDYLINOSITOL -SPECIFIC PHOSPHOLIPASE C; CHAIN: NULL;	PHOSPHATIDYLINOSITOL SPECIFIC
SeqFold Score				86.97		74.76
PMF		-0.20	0.99		0.71	
Verify Score		0.12	0.10		0.09	
PSI BLAST Score		8.1e-11	6.8e-67	6.8e-67	3.4e-34	3.4e-34
End		239	311	312	301	314
Start AA		79	10	01	4	4
Chain ID		ы				
PDB ID		2trc	2plc	2plc	2ptd	2ptd
SEQ ID NO:		604	612	612	612	612

PDB annotation	PHOSPHATIDYLINOSITOL SPECIFIC PHOSPHOLIPASE C	ASE	ASE	STRUCTURAL PROTEIN INTEGRIN- BINDING PROTEIN, INV GENE		NIN HELIX COILED COIL, CONTRACTILE PROTEIN	S; COMPLEX (TRANSCRIPTIONNUCLEARNUCLEA R) NF-AT; TRANSCRIPTION FACTOR, PROTEIN-DNA COMPLEX, NFAT, NF- AT, 2 AP-1, FOS-JUN, QUATERNARY PROTEIN-DNA COMPLEX, CRYSTAL 3 STRUCTURE, TRANSCRIPTION SYNERGY, COMBINATORIAL GENE 4 REGULATION, COMPLEX (TRANSCRIPTION/NUCLEARNUCLEA	C, SERINE PROTEASE SERINE PROTEINASE, TRYPSIN, HYDROLASE	\vdash
Coumpound	CHAIN: NULL;	GLYCOSYLTRANSFERASE CYCLODEXTRIN	GLYCOSYLTRANSFERASE (E.C.2.4.1.19) 1CGT 3	INVASIN; CHAIN: A;	CYCLODEXTRIN GLUCANOTRANSFERASE; CHAIN: A, B;	HUMAN SKELETAL MUSCLE ALPHA-ACTININ 2; CHAIN: A;	NFAT; CHAIN: N; C-FOS; CHAIN: F; C-JUN; CHAIN: J; DNA; CHAIN: A, B;	TRYPSIN; CHAIN: A, B, C, D;	TRYPSIN; CHAIN: A, B, C,
SeqFold Score									210.95
PMF Score		-0.19		-0.19	-0.19	0.09	0.58	1.00	
Verify Score		0.37		0.02	-0.00	0.05	-0.15	1.04	
PSI BLAST Score		2.7e-14		1.4e-31	2.7e-14	2.2e-13	0.0081	1.7e-93	1.7e-93
End AA		339		349	215	201	201	250	250
Start		144		86	75	42	176	23	24
Chain ID				А	∢	A	-	V	<u></u>
PDB CD		lcgt		1cwv	lpam	1quu	1a02	1a0j	1a0j
SEQ NO:		617		617	617	622	627	629	629

SEQ ID NO:	PDB U	Chain 1D	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
629	1a0]	A	24	250	1.4e-73			144.00	BETA-TRYPTASE; CHAIN: A, B, C, D;	SERINE PROTEINASE TRYPSIN-LIKE SERINE PROTEINASE, TETRAMER, HEPARIN. ALLERGY, 2 ASTHMA
629	1ao5	Ą	23	250	5.4e-89			197.00	GLANDULAR KALLIKREIN-13; CHAIN:	SERINE PROTEASE PRORENIN CONVERTING ENZYME (PRECE),
									A, B;	EPIDERMAL GLANDULAR KALLIKREIN, SERINE PROTEASE, PROTEIN MATURATION
629	1ao5	∢	24	250	5.4e-89	1.00	1.00		GLANDULAR KALLIKREIN-13; CHAIN:	SERINE PROTEASE PRORENIN CONVERTING ENZYME (PRECE).
									A, B;	EPIDERMAL GLANDULAR KALLIKREIN, SERINE PROTEASE, PROTEIN MATITRATION
629	laut	ບ	23	249	5.4e-81			138.35	ACTIVATED PROTEIN C;	COMPLEX (BLOOD
									MAI; CHAIN: P;	AUTOPROTHROMBIN IIA;
										HYDROLASE, SERINE PROTEINASE), PLASMA CALCITIM BINDING 2
								-		GLYCOPROTEIN, COMPLEX (BLOOD
629	lazz	A	24	250	3.4e-63		1	144 12	COLL AGENIA SE: CHAIN:	COAGULATION/INFIBITOR)
									A, B; ECOTIN; CHAIN: C,	PROTEASE/INHIBITOR) TRYPSIN
								-	ά	INHIBITOR; COMPLEX (SERINE
										PROTEASE/INHIBITOR), SERINE
							_			PROTEASE, 2 INHIBITOR, COMPLEX, PROTEASE-SUBSTRATE
										INTERACTIONS, 3 COLLAGEN
629	1bio		24	249	2.2e-83			157.25	COMPLEMENT FACTOR D;	SERINE PROTEASE SERINE
		_			_				CHAIN: NULL;	PROTEASE, HYDROLASE,
				_	_					COMPLEMENT, FACTOR D,
										CAIALY IIC 2 IKIAD, SELF-
629	lbqy	A	24	250	5.1e-79			180.19	PLASMINOGEN ACTIVATOR: CHAIN: A B.	BLOOD CLOTTING TSV-PA;
		-	1	1		1	1		ACIIVAIUN, CHAIN, A, D,	FIBRINGL 1313, PLASMINGGEN

PDB annotation	ACTIVATOR, SERINE PROTEINASE, 2 SNAKE VENOM, COMPLEX (HYDROLASE/INHIBITOR), BLOOD CLOTTING	SERINE PROTEASE HYDROLASE, SERINE PROTEASE, DIGESTION, PANCREAS, ZYMOGEN, 2 SIGNAL, MULTIGENE FAMILY	HYDROLASE/HYDROLASE INHIBITOR ENTEROKINASE, HEAVY CHAIN; ENTEROKINASE, LIGHT CHAIN; ENTEROPEPTIDASE, TRYPSINOGEN ACTIVATION, 2 HYDROLASE/HYDROLASE INHIBITOR	SERINE PROTEINASE	COMPLEX (PROTEASE/INHIBITOR) TRYPSIN, COAGULATION FACTOR XA, CHIMERA, PROTEASE, PPACK, 2 CHLOROMETHYLKETONE, COMPLEX (PROTEASE/INHIBITOR)		
Coumpound	GLU-GLY-ARG- CHLOROMETHYLKETONE INHIBITOR; CHAIN: E, F;	TRYPSIN; CHAIN: NULL;	ENTEROPEPTIDASE; CHAIN: A; ENTEROPEPTIDASE; CHAIN: B; VAL-ASP-ASP- ASP-ASP-LYS PEPTIDE; CHAIN: C;	ELASTASE; IELT 4 CHAIN: NULL; IELT 5	COAGULATION FACTOR XA-TRYPSIN CHIMERA; CHAIN: A; D-PHE-PRO- ARG- CHLOROMETHYLKETONE (PPACK) WITH CHAIN: I;	COMPLEX(PROTEINASE/I NHIBITOR) TRYPSIN (E.C.3.4.21.4) COMPLEXED WITH INHIBITOR FROM BITTER IMCT 3 GOURD	COMPLEX(PROTEINASE/I NHIBITOR) TRYPSIN (E.C.3.4.21.4) COMPLEXED WITH INHIBITOR FROM BITTER IMCT 3 GOURD IMCT 4
SeqFold Score		197.58	145.55	140.39	179.71		211.99
PMF Score						1.00	
Verify Score						9.95	
PSI BLAST Score		1.2e-89	2.7e-80	16-67	8.5e-80	6.8e-95	6.8e-95
End AA		250	249	248	250	249	250
Start AA		24	24	23	24	23	24
Chain ID			В		A	A	Y
PDB ID		ldpo	lekb	lelt	1fxy	lmct	Imet
SEQ NO:		629	629	629	629	629	629

PDB annotation		SERINE PROTEINASE SERINE PROTEINASE, GLYCOPROTEIN		 		GROWTH FACTOR 7S NGF; GROWTH FACTOR (BETA-NGF), HYDROLASE - SERINE PROTEINASE 2 (GAMMA-NGF), INACTIVE SERINE PROTEINASE (ALPHA-NGF)	COMPLEX (SERINE PROTEASE/INHIBITOR) TRYPSIN INHIBITOR; SERINE PROTEASE, INHIBITOR, COMPLEX, METAL BINDING SITES, 2 PROTEIN ENGINEERING, PROTEASE. SUBSTRATE INTERACTIONS, 3 METALLOPROTEINS	COMPLEX (SERINE PROTEASE/INHIBITOR) TRYPSIN INHIBITOR; SERINE PROTEASE, INHIBITOR, COMPLEX, METAL BINDING SITES, 2 PROTEIN
Coumpound		NEUROPSIN; CHAIN: A, B;	PLÁSMINOGEN; CHAIN: A, B, C, D;	NERVE GROWTH FACTOR; CHAIN: A, B, G, X, Y, Z;	NERVE GROWTH FACTOR; CHAIN: A, B, G, X, Y, Z;	NERVE GROWTH FACTOR; CHAIN: A, B, G, X, Y, Z;	ECOTIN; CHAIN: A; ANIONIC TRYPSIN; CHAIN: B;	ECOTIN; CHAIN: A; ANIONIC TRYPSIN; CHAIN: B;
SeqFold	Score	235.42	132.25	158.03		203.29		195.91
!	Score				1.00		1.00	
Verify	Score		٠		0.93		0.92	
PSI	BLAST Score	2.7e-88	8.1e-79	8.1e-77	5.4e-91	5.4e-91	6.8e-89	6.8e-89
End	AA	249	250	250	250	250	249	250
Start	AA	24	9	32	24	24	23	24
Chain	ar	Ą	∢	Ą	٥ ا	O	B	В
PDB	a	lnpm	lqrz	lsgf	lsgf	lsgf	Islw	Islw
SEQ	g ö	629	629	629	629	629	629	629

$\overline{}$						
PDB annotation	ENGINEERING, PROTEASE- SUBSTRATE INTERACTIONS, 3 METALLOPROTEINS					
Coumpound		HYDROLASE(SERINE PROTEINASE) TONIN (E.C. NUMBER NOT ASSIGNED) 1TON 4	HYDROLASE(SERINE PROTEINASE) TONIN (E.C. NUMBER NOT ASSIGNED)	HYDROLASE (SERINE PROTEINASE) TRYPSIN (E.C.3.4.21.4) COMPLEXED WITH THE INHIBITOR ITRN 3 DIISOPROPYL-FLUOROPHOSPHOFLUORI DATE (DFP) ITRN 4 HUMAN TRYPSIN, DFP	HYDROLASE (SERINE PROTEINASE) TRYPSIN (E.C.3.4.21.4) COMPLEXED WITH THE INHIBITOR ITRN 3 DIISOPROPYL- FLUOROPHOSPHOFLUORI DATE (DFP) ITRN 4 HUMAN TRYPSIN, DFP	HYDROLASE(SERINE PROTEINASE) TRYPSIN (E.C.3.4.21.4) COMPLEXED WITH BENZAMIDINE INHIBITOR 2TBS 3
SeqFold Score			201.47		198.92	
PMF Score		1.00		1.00		1.00
Verify Score		0.74		0.92		1.09
PSI BLAST Score		1.9e-91	1.9e-91	1.7e-92	1.7e-92	8.5e-91
End		250	250	249	250	250
Start AA		24	24	23	24	23
Chain ID				4	4	
PDB ID		Iton	Iton	Ē	ITT.	2tbs
SEQ ID NO:		629	629	629	629	629

SEQ ID	PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
629	2tbs		24	250	8.5e-91			195.28	HYDROLASE(SERINE PROTEINASE) TRYPSIN (E.C.3.4.21.4) COMPLEXED WITH BENZAMIDINE INHIBITOR 2TBS 3	
629	Sptp		23	250	1.7e-91	1.02	1.00		BETA TRYPSIN; CHAIN: NULL;	SERINE PROTEASE HYDROLASE, SERINE PROTEASE, DIGESTION, PANCREAS, 2 ZYMOGEN, SIGNAL
629	5ptp		24	250	1.7e-91			202.83	BETA TRYPSIN; CHAIN: NULL;	SERINE PROTEASE HYDROLASE, SERINE PROTEASE, DIGESTION, PANCREAS, 2 ZYMOGEN, SIGNAL
632	1a31	្ន	21	170	1.7e-78			157.53	IMMUNOGLOBULIN FAB 13G5; CHAIN: L, H;	IMMUNOGLOBULIN DIELS-ALDER, DISFAVORED REACTION, CATALYTIC ANTIBODY, 2 IMMUNOGLOBULIN
632	1a3r	7	21	691	1.7e-90	0.53	1.00		IGG2A; CHAIN: 1, H; HUMAN RHINOVIRUS CAPSID PROTEIN VP2; CHAIN: P;	COMPLEX (IMMUNOGLOBULIN/VIRAL PEPTIDE) ANTIBODY 8F5; IMMUNOGLOBULIN, ANTIBODY, RHINOVIRUS, NEUTRALIZATION, 2 CONTINUOUS EPITOPE, COMPLEX (IMMUNOGLOBULIN/VIRAL PEPTIDE)
632	1a3r	ı	21	170	1.7e-90			151.40	IGGZA; CHAIN: L, H; HUMAN RHINOVIRUS CAPSID PROTEIN VP2; CHAIN: P;	COMPLEX (IMMUNOGLOBULIN/VIRAL PEPTIDE) ANTIBODY 8FS; IMMUNOGLOBULIN, ANTIBODY, RHINOVIRUS, NEUTRALIZATION, 2 CONTINUOUS EPITOPE, COMPLEX (IMMUNOGLOBULIN/VIRAL PEPTIDE)
632	la4j	1	21	170	1.4e-85			176.88	IMMUNOGLOBULIN, DIELS ALDER CATALYTIC ANTIBODY; CHAIN: L, H, A, B;	IMMUNOGLOBULIN IMMUNOGLOBULIN, ANTIBODY, CATALYTIC ANTIBODY, DIELS ALDER, 2 GERMLINE
632	1ad9	1	21	170	1.7e-88			160.46	FAB FRAGMENT CTM01;	IMMUNOGLOBULIN

		RAGMENT		N.C		30DY FAB',	ACTION		A LIGHT-	(IGEN)		GIOGENIC		NE				-,-	FAB			ion,	FAB	3	lon,	MBRANE		
PDB annotation		ULIN, FAB F	ULIN	UMANISATION	ULIN	ULIN, ANTIE	DOLASE RE	ULIN	ULIN, KAPP. HEADER	TIBODY/AN	COMPLEX	TIGEN), AN	EM	ULIN, IMMU					ULIN CBR96	3ULIN);	ULIN,	ULIN C REG	III IN MBRO	ULIN):	ULIN C'REG	N, TRANSME		
PD		IMMUNOGLOBULIN, FAB FRAGMENT	IMMUNOGLOBULIN BAB	IMMOOCEOBOEIN, FAB FRAGMENT, HUMANISATION	IMMUNOGLOBULIN	IMMUNOGLOBULIN, ANTIBODY FAB,	CATALYST, ALDOLASE REACTION	MMUNOGLOB	IMMUNOGLOBULIN, KAPPA LIGHI- CHAIN DIMER HEADER	COMPLEX (ANTIBODY/ANTIGEN)	FAB-12; VEGF; COMPLEX	(ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	IMMUNE SYSTEM	IMMUNOGLOBULIN, IMMUNE	SYSTEM				IMMUNOGLOBULIN CBR96 FAB	(IMMUNOGLOBULIN);	IMMUNOGLOBULIN	IMMUNOGLOBULIN C REGION,	IMMINOGIORIII IN MBR96 FAB	(IMMUNOGLOBULIN):	IMMUNOGLOBULIN C REGION	GLYCOPROTEIN, TRANSMEMBRANE		
Coumpound		CHAIN: L, H, A, B;	TM01;			IGG2A; CHAIN: L, H;		OBULIN;		Ë		ENDOTHELIAL GROWTH FACTOR: CHAIN: V. W:			HAIN: A,	C; MONOCLONAL	ANTIBODY MRK-16	(HEAVY CHAIN); CHAIN:	IGG FAB (HUMAN IGGI,				IGG FAB (IGG3 KAPPA):				IMMUNOGLOBULIN FABI	EROID
		CHAIN:	ANTIBODY C	CIRCILA	IMMUN	IGG2A; (CHAIN: A, B;	FAB FR	L, H, J, K	FACTOR	MONOCLONAL	ANTIBO	(LIGHT)	C; MON	ANTIBO	(HEAVY	IGG FAB	KAPPA);			IGG FAR	CHAIN: L. H.			FRAGME	ANTI-STEROID
SeqFold Score			164.95		161.73								165.55						175.29				165.08				161.02	
PMF Score								0.0		1.00																		
Verify Score								0.64		29'0																		
PSI BLAST	Score		1e-79		1.7e-84			1.5e-88		1.2e-90			1.7e-85						2.7e-83				5.4e-84				1.1e-83	
End			170		170			170		170			170						170				170	:			170	
Start AA			21		21			7		21	_		21						23				21	:			21	
Chain			1	_	1			∢		<u> </u>			A						L				1	1			L)	
PDB CI			lae6		laxt			1064		16j1			1bln						Icly				1clz				1dbb	
SEQ	Ö		632		632		3	632		632			632						632				632	}			632	

PDB annotation		IMMUNE SYSTEM FAB-IBP COMPLEX CRYSTAL STRUCTURE 2.7A RESOLUTION BINDING 2 OUTSIDE THE ANTIGEN COMBINING SITE SUPERANTIGEN FAB VH3 3 SPECIFICITY	IMMUNOGLOBULIN				CATALYTIC ANTIBODY CATALYTIC ANTIBODY 6D9 CATALYTIC ANTIBODY, ESTER HYDROLYSIS, ESTEROLYTIC, FAB, 2 IMMUNOGLOBULIN	
Coumpound	ANTIBODY 1DBB 3 (IGG1, SUBGROUP 2A, KAPPA 1) COMPLEX WITH PROGESTERONE 1DBB 4	IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B, D, F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H;	4-4-20 (IG*G2A=KAPPA=) FAB FRAGMENT; IFLR 5 CHAIN: L, H; IFLR 6	IMMUNOGLOBULIN FAB FRAGMENT OF HUMANIZED ANTIBODY 4D5, VERSION 4 1FVD 3	IMMUNOGLOBULIN IGG2A FAB FRAGMENT (FAB 17/9) 1HIL 3	IMMUNOGLOBULIN IGG2A FAB FRAGMENT (FAB 17/9) IHIL 3	IMMUNOGLOBULIN 6D9; CHAIN: L, H;	IMMUNOGLOBULIN IGG2A FAB FRAGMENT (FAB 17/9) COMPLEX WITH PEPTIDE OF 11FH 3 INFLUENZA HEMAGGLUTININ HA1
SeqFold Score			169.43			148.78	165.42	
PMF Score		1.00		1.00	1.00			1.00
Verify Score		0.58		0.57	0.45			0.50
PSI BLAST Score		1.7e-91	3.4e-86	3.4e-89	3.4e-90	3.4e-90	1.5e-84	3.4e-90
End AA		170	170	170	169	170	170	169
Start		21	21	21	21	21	21	21
Chain ID		¥	ļ	A	Ą	A	J	J
PDB ID		ldee	i∯.	1fvd	lhil	ıhil	1hyx	lifh
SEQ ID NO:		632	632	632	632	632	632	632

PDB annotation											IMMUNOGLOBULIN	MONOCLONAL ANTIBODY	MONOCLONAL ANTIBODY, FAB- FRAGMENT, REPRODUCTION	MONOCLONAL ANTIBODY	MONOCLONAL ANTIBODY, FAB- FRAGMENT, REPRODUCTION	CATALYTIC ANTIBODY CATALYTIC	ANTIBODY, TRANSITION STATE ANALOGUE	IMMUNE SYSTEM ABZYME, TRANSITION STATE ANALOG,
Coumpound	(STRAIN X47) (RESIDUES 101-107) 1IFH 4	IMMUNOGLOBULIN IGG2A FAB FRAGMENT (FAB 17/9) COMPLEX	WITH PEPTIDE OF 1IFH 3 INFLUENZA	HEMAGGLUTININ HAI (STRAIN X47) (RESIDUES 101-107) 1IFH 4	IMMUNOGLOBULIN IGGI FAB' FRAGMENT (B1312) 11GF 3	IMMUNOGLOBULIN	IMMUNOGLOBULIN FAB FRAGMENT (MC/PC\$603)	IMMUNOGLOBITLIN	IMMUNOGLOBULIN FAB	FRAGMENT (MC/PC\$603) IMCP 4	IGG2A=KAPPA=; 1PLG 4 CHAIN: L, H; 1PLG 5	MONOCLONAL	ANTIBODY 3A2; CHAIN: H, L;	MONOCLONAL	ANTIBODY 3A2; CHAIN: H, L:	IGG2A FAB FRAGMENT	(D2.3); CHAIN: L, H;	IG ANTIBODY D2.3 (LIGHT CHAIN); CHAIN: L; IG
SeqFold Score		148.62			163.94			153.91			165.16			151.88	-	153.63		154.30
PMF Score		_				1.00						0.I						
Verify Score						69.0						0.64						ĺ
PSI BLAST Score		3.4e-90			1.4e-83	1e-93		le-93			3.4e-85	6.8e-95		6.8e-95		1.1e-82		1.1e-82
End		170			170	691		170			170	169		170		170	_,-	170
Start AA		21			21	21		21			21	21		21		21	j	21
Chain ID		1			1	L		1			L	Т		1		1		L
PDB E1		1ifh			ligf	Ітср		Imcp			1plg	1sbs		1sbs		lyec		1yej
SEQ ID NO:		632			632	632		632			632	632		632		632		632

SEQ ID	PDB ID	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
Ö					Score				ANTIBODY D2.3 (HEAVY	IMMUNE SYSTEM
									CHAIN); CHAIN: H;	
632	2fgw	L)	21	170	6.8e-91	0.64	1.00		IMMUNOGLOBULIN FAB	
		-							FRAGMENT OF A	
									HUMANIZED VERSION OF	
									THE ANTI-CD18 2FGW 3	
									ANTIBODY 'H52' (HUH52-	
									OCEAN) ZEOW 4	
3	5		6		200			9		
659	1829	∢	2	81	3.4e-33			51.12	MONOCLONAL	COMPLEX
									ANTIBODI DI.3; CHAIN:	(IMIMUNOCEOBOLIN/RIDROLASE)
									A, B; LISUZIME; CHAIN:	COMPLEX
									<u>ر</u>	(IMIMUNOGLOBOLIN/HIDROLASE),
						-				IMMUNOGLOBULIN V 2 REGION,
										SIGNAL, HIDROLASE,
				-		-				GLYCOSIDASE, BACTERIOLYTIC 3
55			1	:	00,	1				ENCAME, EOG WILLE
cco	18/9	٦		×:	1./e-32			20.86	MUNOCLONAL ANTIBODY D1.3: CHAIN:	IMMUNOGLOBULIN VARIANT
									L, H;	
633	lap2	Ą	20	118	1.7e-33			51.40	MONOCLONAL	IMMUNOGLOBULIN VARIABLE
									ANTIBODY C219; CHAIN:	DOMAIN; SINGLE CHAIN FV,
					-				A, B, C, D;	MONOCLONAL ANTIBODY, C219, P.
						-		-		GLYCOPROTEIN, 2
										IMMUNOGLOBULIN
633	lbd2	ш	22	117	5.1e-38	-0.18	0.99		HLA-A 0201; CHAIN: A;	COMPLEX (MHC/VIRAL
									BETA-2 MICROGLOBULIN;	PEPTIDE/RECEPTOR) HLA A2 HEAVY
									CHAIN: B; TAX PEPTIDE;	CHAIN; COMPLEX (MHC/VIRAL
				-				•	CHAIN: C; T CELL	PEPTIDE/RECEPTOR)
						_		•	RECEPTOR ALPHA;	
				_					CHAIN: D; T CELL	
•								· — ·	RECEPTOR BETA; CHAIN:	
633	Ibec		23	117	3.4e-38	0.14	1.00		14.3.D T CELL ANTIGEN	RECEPTOR T CELL RECEPTOR IBEC

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									RECEPTOR; 1BEC 5 CHAIN: NULL; 1BEC 6	14
633	16fv	L	20	118	8.5e-27			51.90	FV4155; CHAIN: L, H;	IMMUNOGLOBULIN IMMUNOGLOBULIN, FV FRAGMENT, STEROID HORMONE, 2 FINE SPECIFICITY
633	16j1	1	20	117	1.7e-38	-0.21	0.95		FAB FRAGMENT; CHAIN: L, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR; CHAIN: V, W;	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR
633	lbvk	Ą	20	118	6.8e-38			52.24	HULYSII; CHAIN: A, B, D, E; LYSOZYME; CHAIN: C, F;	COMPLEX (HUMANIZED ANTIBODY/HYDROLASE) MURAMIDASE; HUMANIZED ANTIBODY, ANTIBODY COMPLEX, FV, ANTI-LYSOZYME, 2 COMPLEX (HUMANIZED ANTIBODY/HYDROLASE)
633	Idee	4	20	117	6.8e-40	-0.29	0.81		IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B, D, F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H;	IMMUNE SYSTEM FAB-IBP COMPLEX CRYSTAL STRUCTURE 2.7A RESOLUTION BINDING 2 OUTSIDE THE ANTIGEN COMBINING SITE SUPERANTIGEN FAB VH3 3 SPECIFICITY
633	ldfb	J	20	118	3.4e-38	0.22	0.83		IMMUNOGLOBULIN 3D6 FAB 1DFB 3	
633	ıdıf	L	20	117	1.7e-27			52.01	ANTI-DANSYL IMMUNOGLOBULIN IGG2A(S); CHAIN: L, H;	IMMUNOGLOBULIN ANTI-DANSYL FV FRAGMENT FV FRAGMENT, IMMUNOGLOBULIN
633	ldsf	L	20	118	3.4e-25			50.95	ANTICANCER ANTIBODY BI; CHAIN: L, H;	IMMUNOGLOBULIN BIDSFV; MONOCLONAL ANTIBODY, ANTITUMOR, IMMUNOGLOBULIN
633	lfgv	1	20	118	3.4e-40	-0.00	0.99		IMMUNOGLOBULIN FV FRAGMENT OF A HUMANIZED VERSION OF	

PDB annotation																																	
Coumpound		THE ANTI-CD18 IFGV 3 ANTIBODY 'H52' (HUH52- AA FV) IFGV 4	IMMUNOGLOBULIN FV	FRAGMENT OF A	HUMANIZED VERSION OF	THE ANTI-CD18 1FGV 3	ANTIBODY 'H52' (HUH52-	AA FV) IFGV 4	IMMUNOGLOBULIN FV	FRAGMENT OF	HUMANIZED ANTIBODY	MAKENDOCI OBITE IN EV	IMIMONOGLOBOLIN FV	FRAGMENI OF	HUMANIZED ANTIBODY	TO THE STORY OF THE PUBLICATION	IMMUNOGLOBULIN FAB	FRAGMENT OF	HUMANIZED ANTIBODY	4D5, VERSION 4 1FVD 3	IMMUNOGLOBULIN	IMMUNOGLOBULIN VL	DOMAIN (VARIABLE	DOMAIN OF KAPPA LIGHT	IIVL 3 CHAIN) OF	DESIGNED ANTIBODY	M29B 1IVL 4	IMMUNOGLOBULIN	MURINE ANTIBODY 26-10	VL DOMAIN (NMR, 15	MAATA STRICTIBES	IMAS STRUCTORES)	TIMIN +
SeqFold Score			54.70									50 10	01.26			1					52.59							54.30					
PMTF Score									98.0							3	 88.																
Verify Score									-0.06							000	0.08						_								_		
PSI BLAST	Score		3.4e-40			_			3.4e-40			2 40 40	3.46-40			;	6.8e-41				3.4e-28		_	-		-		8.5e-27					
End AA			118						118			1,0	011				× 1.				1117							118					
Start AA			20						20			20	3			8	70				70							20					
Chain ID			7						A			<	۲				∢				Ą		_	_									
PDB ID			lfgv						1fvc			16,0	3			1 9	IVG				livi							lmaj					
SEQ ID	NO:		633				_		633			633	ŝ			5	250				633				_			633					

PDB annotation	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL			IMMUNOGLOBULIN	OXYGEN TRANSPORT OXYGEN TRANSPORT, HEME, RESPIRATORY PROTEIN, ERYTHROCYTE	OXYGEN TRANSPORT OXYGEN TRANSPORT	-
Coumpound	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	IMMUNOGLOBULIN FAB FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4	IMUNOGLOBULIN IMMUNOGLOBULIN VL DOMAIN (VARIABLE DOMAIN OF KAPPA ZIMN 3 LIGHT CHAIN) OF MCPC603 MUTANT IN WHICH ZIMN 4 COMPLEMENTARITY- DETERMINING REGION I HAS BEEN REPLACED BY ZIMN 5 THAT FROM MOPC167 ZIMN 6	IMMUNOGLOBULIN (LIGHT CHAIN); CHAIN: A, C, E, G; IMMUNOGLOBULIN (HEAVY CHAIN); CHAIN: B, D, F, H;	HEMOGLOBIN; CHAIN: A, B	HEMOGLOBIN; CHAIN: A, E, C, F;	OXYGEN TRANSPORT
SeqFold Score			50.13	52.37			
PMF Score	1.00	0.43			1.00	1.00	1.00
Verify Score	0.00	-0.09			-0.66	-0.68	-0.71
PSI BLAST Score	1.7e-38	6.8e-40	5.1e-34	1.7e-33	1.7e-35		1.7e-36
End	115	118	118	118	77	77	72
Start	20	20	20	20	_		_
Chain 10	В	1		A	В	ப	В
PDB CI	Itcr	2fgw	2imn	43c9	1a4f		1bab
Se Se Se	633	633	633	633	635		635

PDB annotation		OXYGEN TRANSPORT OXYGEN TRANSPORT, CHIMERA PROTEIN, RESPIRATORY PROTEIN, HEME		OXYGEN STORAGE/TRANSPORT HB D; HB D HEMOGLOBIN D (R-STATE) 1, HEMOGLOBIN, AVIAN, HIGH 2 COOPERATIITY, OXYGEN TRANSPORT			OXYGEN TRANSPORT HEME, OXYGEN TRANSPORT, RESPIRATORY PROTEIN, ERYTHROCYTE	OXYGEN TRANSPORT X-RAY STUDY, PORCINE HEMOGLOBIN, ARTIFICIAL HUMAN BLOOD, 2 OXYGEN TRANSPORT
Coumpound	HEMOGLOBIN THIONVILLE ALPHA CHAIN MUTANT WITH VAL 1 IBAB 3 REPLACED BY GLU AND AN ACETYLATED MET BOUND TO THE IBAB 4 AMINO TERMINUS IBAB 5	MODULE-SUBSTITUTED CHIMERA HEMOGLOBIN BETA-ALPHA; CHAIN: A, B, C, D;	OXYGEN TRANSPORT HEMOGLOBIN (DEOXY, HUMAN FETAL F=/II\$=) IFDHG 1 IFDHH 2	HEMOGLOBIN D; CHAIN: A, C; HEMOGLOBIN D; CHAIN: B, D;	OXYGEN TRANSPORT HEMOGLOBIN (DEOXY) IHDA 3	OXYGEN TRANSPORT HEMOGLOBIN (SICKLE CELL) 1HDS 4	HEMOGLOBIN (DEOXY); CHAIN: A, B;	PORICINE HEMOGLOBIN (ALPHA SUBUNIT); CHAIN: A, C; PORICINE HEMOGLOBIN (BETA
SeqFold Score								
PMF Score		1.00	1.00	1.00	66'0	0.93	1.00	1.00
Verify Score		-0.68	-0.70	-0.76	-0.57	-0.73	-0.46	-0.64
PSI BLAST Score		5.1e-35	1.7e-38	1e-35	1.7e-33	6.8e-31	1.2e-36	1.5e-34
End AA		75	77	77	77	11	11	77
Start AA		_		-	-	_	_	_
Chain ID		۲	ى ن	В	В	æ	В	a
PDB CD		1ch4	1fdh	1hbr	1hda	Ihds	libe	Ідрм
SEQ NO:		635	635	635	635	635	635	635

PDB annotation	((IMMUNOGLOBULIN/HYDROLASE), IMMUNOGLOBULIN V 2 REGION,	SIGNAL, HYDROLASE, GLYCOSIDASE, BACTERIOLYTIC 3 ENZYME, EGG WHITE		IMMUNOGLOBULIN	IN: IMMUNOGLOBULIN, FAB FRAGMENT	╀	AIN: (IMMUNOGLOBULIN/AUTOANTIGEN)	(IMMUNOGLOBULIN/AUTOANTIGEN),	KHEUMATUID FACTOR 2 AUTO- ANTIBODY COMPLEX	H; IMMUNOGLOBULIN HUMAN FAB,	AFFINITY, CRYSTAL 2 PACKING	MOTE, PROGRAMMING PROPENSITY	IOCKISIALLIZE, 3 IMMUNOGLOBULIN				(OXIDOREDUCTASE/ANTIBODY),
Coumpound	SUBUNIT); CHAIN: B, D		MONOCLONAL ANTIBODY D1.3; CHAIN: A, B; LYSOZYME; CHAIN:	ပ်		MONOCLONAL ANTIBODY DI.3; CHAIN: I H:	FAB FRAGMENT,	ANTIBODY ASB7; CHAIN: A, B, C, D;	IGG4 REA; CHAIN: A; RF-	AN IGM/LAMBDA; CHAIN:	î î		FAB B7-15A2; CHAIN: L, H;				CYTOCHROME C	ANTIBODY FV	FRAGMENT; CHAIN: C, D;	
SeqFold Score			26.90			57.44	51.63										52.73			
PMF Score									0.81				0.74							
Verify Score									0.19				0.05							
PSI BLAST Score			1.7e-37			le-35	6.8e-56		1.5e-76				1.2e-68				1.7e-33			
End		į	131			131	200		189				188				131			
Start AA			19			61	5		21				21				61			
Chain ID			V			7	A		1				7				Ω			
PDB CI			1a2y			1a7q	lad0		ladq				laqk				larl			
SEQ NO:			637			637	637		637				637				637			

PDB annotation	TRANSMEMBRANE, CYTOCHROME OXIDASE, ANTIBODY COMPLEX	IMMUNE SYSTEM BENCE-JONES; IMMUNOGLOBULIN, AMYLOID, IMMUNE SYSTEM	IMMUNOGLOBULIN IMMUNOGLOBULIN, KAPPA LIGHT- CHAIN DIMER HEADER	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA A2 HEAVY CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR)	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	IMMUNOGLOBULIN BENCE-JONES PROTEIN; IBJM 8 BENCE JONES, ANTIBODY, MULTIPLE QUATERNARY STRUCTURES IBJM 13	COMPLEX (HUMANIZED ANTIBODY/HYDROLASE) MURAMIDASE; HUMANIZED ANTIBODY, ANTIBODY COMPLEX, FV, ANTI-LYSOZYME, 2 COMPLEX (HUMANIZED
Coumpound		BENCE-JONES KAPPA I PROTEIN BRE; CHAIN: A, B, C;	IMMUNOGLOBULIN; CHAIN: A, B;	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	FAB FRAGMENT; CHAIN: L, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR: CHAIN: V. W:	FAB FRAGMENT; CHAIN: L, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR: CHAIN: V. W:	LOC - LAMBDA I TYPE LIGHT-CHAIN DIMER; IBJM 6 CHAIN: A, B; IBJM	HULYSI I; CHAIN: A, B, D, E; LYSOZYME; CHAIN: C, F;
SeqFold Score		55.76	50.36	51.40		52.82		54.69
PMF Score					0.64		0.53	
Verify Score					0.21		0.05	
PSI BLAST Score		1.4e-39	3.4e-55	1.6e-09	1.7e-57	1.7e-57	8.5e-67	3.4c-41
End AA		131	200	199	921	200	188	131
Start AA		61	19	20	61	61	21	19
Chain 10		A	A	Q	<u>.</u>	7	Ą	¥
PDB UD		1b0w	1b6d	1bd2	16j1	16j1	1bjm	15vk
SEQ ID NO:		637	637	637	637	637	637	637

	_									_																
PDB annotation		ANTIBODY/HYDROLASE)	IMMUNE SYSTEM REIV, STABILIZED IMMUNOGLOBULN FRAGMENT, BENCE-JONES 2 PROTEIN, IMMUNE SYSTEM	ANTIBODY THERAPEUTIC, ANTIBODY. CD52											-		COMPLEX (ANTIBODY/ANTIGEN)	CYTOKINE RECEPTOR, COMPLEX	(ANTIBODY/ANTIGEN), 2	TRANSMEMBRANE, GLYCOPROTEIN	IMMUNOGLOBULIN	IMMUNOGLOBULIN, BENCE JONES	PROTEIN			
Coumpound			IG KAPPA CHAIN V-I REGION REI; CHAIN: A, B;	CAMPATH-IH:LIGHT CHAIN; CHAIN: L;	CAMPATH-1H:HEAVY	CHAIN; CHAIN: H;	CHAIN: P;	IMMUNOGLOBULIN 3D6 FAB IDFB 3	IMMUNOGLOBULIN FV	FRAGMENT OF A	HUMANIZED VERSION OF	THE ANTI-CD18 1FGV 3	ANTIBODY 'H52' (HUH52-	AA FV) 1FGV 4	IMMUNOGLOBULIN M	(IG-M) FV FRAGMENT IIGM 3	ANTIBODY A6; CHAIN: L,	H; INTERFERON-GAMMA	RECEPTOR ALPHA CHAIN;	CHAIN: I;	LAMBDA III BENCE JONES	PROTEIN CLE; CHAIN: A,	Б	IMMUNOGLOBULIN	HETEROLOGOUS LIGHT	CHAIN DIMER 1MCW 3
SeqFold Score			56.65	50.97				54.95	57.59						55.75		56.16									
PMF Score																					0.43		100	-0.05	·	
Verify Score																					0.12			0.00	_	
PSI BLAST	Score		1.7e-41	1.2e-53				1.4e-54	3.4e-43						6.8e-42		3.4e-43			$\overline{}$	1.2e-70			1.7e-59		
End			132	200				200	131						140		161				189		12.0	<u> </u>		
Start AA			17	19				61	61						6		19				21		;			
Chain ID			4	J				T	7					ļ	1		7				∀		١	*		
PDB ID			lbww	lce!				1dfb	Ifgv					:	ligm		1j.j.							ımcw		
SEQ	ÿ		637	637				637	637					3	رج ا		637				637		637	è		

PDB annotation		IMMUNOGLOBULIN.	IMMUNE SYSTEM HUMAN TCR/PEPTIDE/MHC COMPI RX HI A.	A2, HTLV-1, TAX, TCR, T 2 CELL	RECEPTOR, IMMUNE SYSTEM			RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN,	SIGNAL																	
Coumpound	(/MCG\$-/WEIR\$ HYBRID)	NIG9 (IGG1=LAMBDA=); CHAIN: L, H;	MHC CLASS I HLA-A; CHAIN: A: BETA-2	MICROGLOBULIN; CHAIN:	B; TAX PEPTIDE P6A;	CHAIN: C; HIMAN 1-CELL RECEPTOR; CHAIN: D;	HLA-A 0201; CHAIN: E;	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;		IMMUNOGLOBULIN WAT,	A VARIABLE DUMAIN	FKUM	IMMUNOGLOBOLIN	(BENCE-JONES PROTEIN)	1WTL 4	IMMUNOGLOBULIN	IMMUNOGLOBULIN FAB 2FB4 4	IMMUNOGLOBULIN FAB	FRAGMENT OF A	HUMANIZED VERSION OF	THE ANTI-CD18 2FGW 3	ANTIBODY 'H52' (HUH52-	OZ FAB) 2FGW 4	IMMUNOGLOBULIN	I AMBDA I IGHT CHAIN	DIMER (/MCG\$) 2MCG 3
SeqFold Score		51.23	52.06					52.17		55.36								52.58								
PMF Score																0.47								0.31		
Verify Score					-	-						_				0.14								0.12		
PSI BLAST	Score	1.5e-21	1.4e-12					6.8e-17		8.5e-41						1.2e-68		3.4e-57						5.1e-68		
End AA		200	200					700		131						188		200						188		
Start AA		19	20					20		6						20		61			_			21		\neg
Chain		Н	Q					∢		V						<u> </u>		1							_	
PDB CI		lngp	lqm					Itcr		Iwt						2fb4		2fgw						2mcg		
SEQ	Ö	637	637					637		637						637		637						637		

	1		 	\neg	Τ	<u> </u>		Τ			•		Γ.	
PDB annotation					COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING PROTEIN		COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN		COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING PROTEIN		COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING PROTEIN
Coumpound	(TRIGONAL FORM) 2MCG 4	IMMUNOGLOBULIN BENCE-*JONES PROTEIN (LAMBDA, VARIABLE DOMAIN) 2RHE 4	IMMUNOGLOBULIN FAB FRAGMENT FROM HUMAN IMMUNOGLOBULIN IGGI	(LAMBDA, HIL) 8FAB 3	QGSR ZINC FINGER PEPTIDE; CHAIN: A;	DUPLEX OLIGONUCLEOTIDE	BINDING SITE; CHAIN: B, C;	QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX	OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C;	QGSR ZINC FINGER PEPTIDE; CHAIN: A;	DUPLEX OLIGONUCLEOTIDE	BINDING SITE; CHAIN: B, C;	QGSR ZINC FINGER PEPTIDE: CHAIN: A:	DUPLEX, OLIGONUCLEOTIDE
SeqFold Score		55.78			59.24						,-,-			
PMF Score			0.87					0.13		1.00			1.00	
Verify Score			0.10					0.29		0.32			-0.17	
PSI BLAST Score		le-47	1.7e-69		1.2e-29			1.2e-29		1.2e-27			5.4e-29	
End		138	189		84			110		82			84	
Start AA		20	23		-			28		2			2	
Chain ID			4		A			A		4			¥.	
PDB ID		2rhe	8fab		lalh			lalh		lalh			lalh	
SEQ NO NO		637	637		639			639		639			639	

PDB annotation		COMPLEX (ZINC FINGER/DNA) ZINC FINGER PROTEIN, DNA	INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	INTERACTION, PROTEIN DESIGN, 2 CRASTAL STRICTLINE COMPLEX	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	INTERACTION, PROTEIN DESIGN, 2	CRISIAL SIRUCIORE, COMPLEX (ZINC FINGER/DNA)	COMPLEX (TRANSCRIPTION	REGULATION/DNA) COMPLEX	(TRANSCRIPTION	REGULATION/DNA), RNA	POLYMERASE III, 2 TRANSCRIPTION	INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION PEGIT ATTOMONA) VALGE VANG 1:	TRANSCRIPTION INITIATION	INITIATOR ELEMENT, YY1, ZINC 2	FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX
Coumpound	BINDING SITE; CHAIN: B, C;	DNA; CHAIN: A, B, D, E;	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;		TFIIIA; CHAIN: A, D; 5S	RIBOSOMAL RNA GENE;	CHAIN: B, C, E, F;				YYI; CHAIN: C; ADENO-	NITIATOR ELEMENT	DNA; CHAIN: A, B;	
SeqFold Score		83.70																			3	28.78			
PMF Score					0.71			1.00				1.00				0.01				_					•
Verify Score					0.14			0.07				0.44				-0.26	•								
PSI BLAST Score		3.4e-49			3.4e-49			6.8e-48				3.4e-14				5.1e-28					;	3.46-32			
End		83			110			82				82				112					1	=			
Start		-			27			2				55				2			•		1.	_			
Chain		ပ			၁			ပ				g				A	·				,	<u>۔</u> ن			
PDB ID		Imey			lmey			lmey				lmey				1tf6						Inpd			
SEQ ID		639			639			639				639				639						659			

PDB annotation	(TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1;	TRANSCRIPTION INITIATION,	INITIATOR ELEMENT, YY1, ZINC 2	FINGER PROTEIN, DNA-PROTEIN	RECOGNITION, 3 COMPLEX	(TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION	REGULATION/DNA) YING-YANG 1;	TRANSCRIPTION INITIATION,	INITIATOR ELEMENT, YY1, ZINC 2	FINGER PROTEIN, DNA-PROTEIN	RECOGNITION, 3 COMPLEX	(TRANSCRIPTION REGULATION/DNA)	TRANSCRIPTION REGULATION	TRANSCRIPTION REGULATION,	ADR1, ZINC FINGER, NMR	COMPLEX (DNA-BINDING	PROTEIN/DNA) FIVE-FINGER GLI; GLI,	ZINC FINGER, COMPLEX (DNA-	BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING	PROTEIN/DNA) FIVE-FINGER GLI; GLI,	ZINC FINGER, COMPLEX (DNA-	BINDING PROTEIN/DNA)	COMPLEX (INHIBITOR/NUCLEASE)	COMPLEX (INHIBITOR/NUCLEASE),	COMPLEX (RI-ANG), HYDROLASE 2	MOLECULAR RECOGNITION, EPITOPE	MAPPING, LEUCINE-RICH 3 REPEATS	COMPLEX (NUCLEAR PROTEINRNA)
Coumpound		YY1; CHAIN: C; ADENO- ASSOCIATED VIRUS P5	INITIATOR ELEMENT	DNA; CHAIN: A, B;				YY1; CHAIN: C; ADENO-	ASSOCIATED VIRUS PS	INITIATOR ELEMENT	DNA; CHAIN: A, B;				ADRI; CHAIN: NULL;			ZINC FINGER PROTEIN	GLII; CHAIN: A; DNA;	CHAIN: C, D;	The state of the s	ZINC FINGER PROTEIN	GLII; CHAIN: A; DNA;	CHAIN: C, D;	1	RIBONUCLEASE	INHIBITOR; CHAIN: A, D;	ANGIOGENIN; CHAIN: B,	ű		U2 RNA HAIRPIN IV; CHAIN: O. R: U2 A':
SeqFold Score															51.76											-					
PMF Score		0.95						0.24										-0.03				0.18				0.36					0.63
Verify Score		0.09						-0.02										0.13				0.02				-0.17					0.14
PSI BLAST Score		1.4e-28						3.4e-32							le-15			5.1e-29				5.4e-24				5.4e-15					1.6e-15
End		82						9							68			109				82				181			•		152
Start		1						7							59			17				4				25					40
Chain ID		၁						ပ										∢				∀				A					Ψ
PDB ID		lubd						1npq							2adr			2gli				2gli				la4y			_		1a9n
SEQ ID NO:		639						639							639			639				639				640					640

PDB m	Chain	Start	End	ISA TA	Verify	PMF	SeqFold	Coumpound	PDB annotation
. 1		5	VV.	Score	31016	2006	Score		
								CHAIN: A, C; U2 B"; CHAIN: B, D;	RNA, SNRNP, RIBONUCLEOPROTEIN
la9n	A	28	182	2.7e-22	0.50	0.78		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
								CHAIN: Q, K; 02 A; CHAIN: A. C: U2 B";	COMPLEA (NOCLEAR PROTEIN/RNA), RNA. SNRNP.RIBONIICI.EOPROTEIN
١								CHAIN: B, D;	
la9n	A	82	203	2.2e-17	0.00	-0.09		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
								CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
								CHAIN: A, C; U2 B"; CHAIN: B, D;	RNA, SNRNP, RIBONUCLEOPROTEIN
1a9n	၁	40	159	1.4e-15	0.24	0.88		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
								CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
								CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
1								CHAIN: B, D;	
la9n	ပ	28	182	1.4e-22	0.47	69.0		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
								CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
								CHAIN: A, C; U2 B";	RNA, SNRNP, RIBONUCLEOPROTEIN
١								CHAIN: B, D;	
1a9n	ပ	82	203	1.4e-17	0.55	0.15		U2 RNA HAIRPIN IV;	COMPLEX (NUCLEAR PROTEIN/RNA)
								CHAIN: Q, R; U2 A';	COMPLEX (NUCLEAR PROTEIN/RNA),
								CHAIN: A, C; U2 B"; CHAIN: B. D:	RNA, SNRNP, RIBONUCLEOPROTEIN
1086	A	248	374	5.4e-09	-0.18	90.0		AXONIN-1; CHAIN: A;	CELL ADHESION NEURAL CELL
lcs6	A	334	426	1.7e-08	0.10	-0.14		AXONIN-1: CHAIN: A:	CELL ADHESION NEURAL CELL
									ADHESION
1d0b	A	56	204	1.7e-22	-0.20	0.01		INTERNALIN B; CHAIN: A;	CELL ADHESION LEUCINE RICH
								1	REPEAT, CALCIUM BINDING, CELL ADHESION
1d0b	¥	19	256	6.8e-22	0.18	0.09		INTERNALIN B; CHAIN: A;	CELL ADHESION LEUCINE RICH
					-			1	REPEAT, CALCIUM BINDING, CELL ADHESION
1dce	A	24	E	8.5e-10	0.22	0.72		RAB	TRANSFERASE CRYSTAL
ı				⊣				GERANYLGERANYLTRAN	STRUCTURE, RAB

PDB annotation	GERANYLGERANYLTRANSFERASE, 2.0 A 2 RESOLUTION, N- FORMYLMETHIONINE, ALPHA SUBUNIT, BETA SUBUNIT	TRANSFERASE CRYSTAL STRUCTURE, RAB GERANYLGERANYLTRANSFERASE, 2.0 A 2 RESOLUTION, N- FORMYLMETHIONINE, ALPHA SUBUNIT, BETA SUBUNIT	CONTRACTILE PROTEIN LEUCINE- RICH REPEAT, BETA-BETA-ALPHA CYLINDER, DYNEIN, 2 CHLAMYDOMONAS, FLAGELLA	CONTRACTILE PROTEIN LEUCINE- RICH REPEAT, BETA-BETA-ALPHA CYLINDER, DYNEIN, 2 CHLAMYDOMONAS, FLAGELLA	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF2; FGFR2; IMMUNOGLOBULIN (IG)LIKE DOMAINS BELONGING TO THE 1-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS, B-TREFOIL FOLD	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF1; FGFR1; IMMUNOGLOBULIN (IG) LIKE DOMAINS BELONGING TO THE 1-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS, B-TREFOIL FOLD
Coumpound	SFERASE ALPHA SUBUNIT; CHAIN: A, C; RAB GERANYLGERANYLTRAN SFERASE BETA SUBUNIT; CHAIN: B, D;	RAB GERANYLGERANYLTRAN SFERASE ALPHA SUBUNIT; CHAIN: A, C; RAB GERANYLGERANYLTRAN SFERASE BETA SUBUNIT; CHAIN: B, D;	OUTER ARM DYNEIN; CHAIN: A;	OUTER ARM DYNEIN; CHAIN: A;	FIBROBLAST GROWTH FACTOR 2; CHAIN: A, B, C, D; FIBROBLAST GROWTH FACTOR RECEPTOR 2; CHAIN: E, F, G, H;	FIBROBLAST GROWTH FACTOR 1; CHAIN: A, B; FIBROBLAST GROWTH FACTOR RECEPTOR 1; CHAIN: C, D;
SeqFold Score						
PMF Score		0.84	0.17	0.23	0.05	0.05
Verify Score		-0.04	-0.33	-0.20	-0.76	-0.73
PSI BLAST Score		1.7e-09	3.4e-12	1.ie-15	1.7e-05	1.7e-05
End		183	135	182	372	372
Start AA	 	22	20	75	340	340
Chain ID		V	∢	Ą	ជា	၁
PDB ID		Idce	1ds9	1ds9	lev2	levt
SEQ ID NO:		640	640	640	640	640

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
640	lfna		615	569	0.0027	0.08	0.35		CELL ADHESION PROTEIN FIBRONECTIN CELL- ADHESION MODULE TYPE III-10 1FNA 3	
640	1601	A	45	111	1.7e-06	-0.15	0.11		NUCLEAR RNA EXPORT FACTOR 1; CHAIN: A, B;	RNA BINDING PROTEIN TAP (NFX1); RIBONUCLEOPROTEIN (RNP,RBD OR RRM) AND LEUCINE-RICH-REPEAT 2 (LRR)
640	1f01	В	45	111	1.7e-06	-0.46	0.11		NUCLEAR RNA EXPORT FACTOR 1; CHAIN: A, B;	RNA BINDING PROTEIN TAP (NFX1); RIBONUCLEOPROTEIN (RNP,RBD OR RRM) AND LEUCINE-RICH-REPEAT 2 (LRR)
640	Ifqv	A	52	192	1.46-10	0.01	-0.03		SKP2; CHAIN: A, C, E, G, I, K, M, O; SKP1; CHAIN: B, D, F, H, J, L, N, P;	LIGASE CYCLIN A/CDK2- ASSOCIATED PROTEIN P45; CYCLIN A/CDK2-ASSOCIATED PROTEIN P19; SKP1, SKP2, F-BOX, LRR, LEUCINE- RICH REPEAT, SCF, UBIQUITIN, 2 E3, UBIQUITIN PROTEIN LIGASE
640	162	A	52 .	181	8.1e-16	0.02	0.23		SKP2; CHAIN: A, C; SKP1; CHAIN: B, D;	LIGASE CYCLIN A/CDK2- ASSOCIATED P45; CYCLIN A/CDK2- ASSOCIATED P19; SKP1, SKP2, F-BOX, LRRS, LEUCINE-RICH REPEATS, SCF, 2 UBIQUITIN, E3, UBIQUITIN PROTEIN LIGASE
640	2bnh		52	183	1.6e-19	-0.09	0.29		RIBONUCLEASE INHIBITOR; CHAIN: NULL;	ACETYLATION RNASE INHIBITOR, RIBONUCLEASE/ANGIOGENIN INHIBITOR ACETYLATION, LEUCINE- RICH REPEATS
641	lalh	4	273	348	1.1e-21	-0.09	0.95		QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B,	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN

PDB UD	Chain ID	1 Start AA	End AA	PSI BLAST Score	Verify Score	PMF	SeqFold Score	Coumpound	PDB annotation
		-		2030				Ö	
lath	4	296	376	3.4e-31	-0.27	0.99		QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B,	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN
lalh	V V	536	379	3.4e-31			72.92	CS. QGSR ZINC FINGER PEPTIDE; CHAIN: A; DUPLEX OLIGONUCLEOTIDE BINDING SITE; CHAIN: B, C.	COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC FINGER, DNA-BINDING PROTEIN
1buo	4	w	127	1e-20			58.53	PROMYBLOCYTIC LEUKEMIA ZINC FINGER PROTEIN PLZF; CHAIN: A;	GENE REGULATION POZ DOMAIN; PROTEIN-PROTEIN INTERACTION DOMAIN, TRANSCRPTIONAL 2 REPRESSOR, ZINC-FINGER PROTEIN, X-RAY CRYSTALLOGRAPHY, 3 PROTEIN STRUCTURE, PROMYELOCYTIC LEUKEMIA, GENE REGULATION
onql	∢	9	119	1e-20	G.	1.00		PROMYELOCYTIC LEUKEMIA ZINC FINGER PROTEIN PLZF; CHAIN: A;	GENE REGULATION POZ DOMAIN; PROTEIN-PROTEIN INTERACTION DOMAIN, TRANSCRIPTIONAL 2 REPRESSOR, ZINC-FINGER PROTEIN, X-RAY CRYSTALLOGRAPHY, 3 PROTEIN STRUCTURE, PROMYELOCYTIC LEUKEMIA, GENE REGULATION
lmey	O 20	247	320	5.1e-37	0.03	0.21		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)

SEQ ID	PDB ID	Chain ID	Start AA	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
NO:	Imey	U	273	348	Score 2.7e-23	-0.22	0.94		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX
641	Imey	U	295	377	3.4e-51			78.38	DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	(ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX
641	lmey	U	296	376	3.4e-51	0.14	0.99		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER DNA)
641	1mey	ပ	323	422	1.7e-47	-0.36	0.17		DNA; CHAIN: A, B, D, E; CONSENSUS ZINC FINGER PROTEIN; CHAIN: C, F, G;	COMPLEX (ZINC FINGER/DNA) ZINC FINGER, PROTEIN-DNA INTERACTION, PROTEIN DESIGN, 2 CRYSTAL STRUCTURE, COMPLEX (ZINC FINGER/DNA)
641	<u> </u>	∢	295	380	5.1e-19			59.21	TRANSCRIPTION FACTOR IIIA; CHAIN: A; 5S RNA GENE; CHAIN: E, F;	COMPLEX (TRANSCRIPTION REGULATION/DNA) TFIIIA; 5S GENE; NMR, TFIIIA, PROTEIN, DNA, TRANSCRIPTION FACTOR, 5S RNA 2 GENE, DNA BINDING PROTEIN, ZINC FINGER, COMPLEX 3
641	1tf6	⋖	271	424	3.4e-32	-0.14	0.12		TFIIIA; CHAIN: A, D; SS RIBOSOMAL RNA GENE; CHAIN: B, C, E, F;	COMPLEX (TRANSCRIPTION REGULATION/DNA) COMPLEX (TRANSCRIPTION REGULATION/DNA), RNA POLYMERASE III, 2 TRANSCRIPTION INITIATION, ZINC FINGER PROTEIN
641	1tf6	A	295	440	3.4e-32			86.69	TFIIIA; CHAIN: A, D; 5S	COMPLEX (TRANSCRIPTION

Verify PMF SeqFold Coumpound PDB annotation ST Score Score	CHAIN: B, C, E, F; (TRANSCRIPTION CHAIN: B, C, E, F; REGULATION/DNA), RNA POLYMERASE III, 2 TRANSCRIPTION INITIATION, ZINC FINGER, PROTEIN	87.61 YY1; CHAIN: C; ADENO-ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	-0.29 0.89 YYI; CHAIN: C; ADENO. ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	77.15 ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; I CHAIN: C, D;	1 -0.27 0.40 ZINC FINGER PROTEIN COMPLEX (DNA-BINDING GLII; CHAIN: A; DNA; PROTEIN/DNA) FIVE-FINGER GLI; GLII; CHAIN: C, D; ZINC FINGER, COMPLEX (DNA-BINDING PROTEIN/DNA)	2 51.26 NEURAMINIDASE; CHAIN: COMPLEX (ANTIBODY/ANTIGEN) N; SINGLE CHAIN COMPLEX (ANTIBODY/ANTIGEN), ANTIBODY; CHAIN: H, L; SINGLE-CHAIN ANTIBODY, 2 GI VOOSY ATED PROTEIN	
PSI Ver BLAST Scc Score		5.1e-35	5.1e-35 -0.2	8.5e-31	8.5e-31 -0.2	3.4e-42	
End		377	376	378	378	126	
Start		273	278	233	246	20	1
Chain ID		U	ပ	A	∀	1	
PDB ID		Jubd	lubd	2gli	2gli	1a14	
SEQ ID NO:		641	641	641	641	646	!

_		_	T		1			1						_																
PDB annotation		IMMUNE SYSTEM	IMMUNOGLOBULIN	IMMUNOGLOBULIN, KAPPA LIGHT- CHAIN DIMER HEADER	COMPLEX (ANTIBODY/ANTIGEN)	FAB-12; VEGF; COMPLEX	(ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	COMPLEX (HUMANIZED	ANTIBODY/HYDROLASE)	MURAMIDASE; HUMANIZED	ANTIBODY, ANTIBODY COMPLEX,	FV, ANTI-LYSOZYME, 2 COMPLEX	(HUMANIZED	IMMUNE SYSTEM REIV. STABILIZED	IMMUNOGLOBULIN FRAGMENT,	BENCE-JONES 2 PROTEIN, IMMUNE	SYSIEM	IMMUNE SYSTEM REIV, STABILIZED IMMI NOGLOBLIL IN FRAGMENT	BENCE-JONES 2 PROTEIN, IMMUNE	SYSTEM	ANTIBODY THERAPEUTIC,	ANTIBODY, CD52					IMMUNE SYSTEM FAB-IBP COMPLEX	CRYSTAL STRUCTURE 2.7A	KESOLUTION BINDING 2 OUTSIDE	SHERANTIGEN COMBINING SHE SHERANTIGEN FAR VH3 3
Coumpound		B, C;	IMMUNOGLOBULIN;	CHAIN: A, B;	FAB FRAGMENT; CHAIN:	L, H, J, K; VASCULAR	ENDOTHELIAL GROWTH FACTOR; CHAIN: V, W;	HULYS11; CHAIN: A, B, D,	E; LYSOZYME; CHAIN: C,	÷				IG KAPPA CHAIN V.1	REGION REI; CHAIN: A, B;			IG KAPPA CHAIN V.1 REGION REI: CHAIN: A. B:			CAMPATH-1H:LIGHT	CHAIN; CHAIN: L;	CAMPATH-IH:HEAVY	CHAIN; CHAIN: H;	PEPTIDE ANTIGEN;	CHAIN: P;	IGM RF 2A2; CHAIN: A, C,	E; IGM RF 2A2; CHAIN: B,	C PRIORIC BROTTER A	CHAIN: G. H.
SeqFold	Score							54.02						51.17													_			
PMF	alose		1.00		1.00												5	3			96.0						00.			
Verify	2000		0.21		0.35												5				0.37						0.18			
PSI RI AST	Score		1.7e-51		1.7e-53			1.7e-49		_				le-51			12.61	16-91		-	5.1e-50					-	1.2e-54			
End	5		126		126			130						129			127	77			126					3	971			
Start			20		50			70						28			ç	2		7	702					1	7			
Chain TD			٧		7			∢						V			-	ς		,						1				
PDB			1 poq		lbj1			lbvk _						Jbww			1811111	*	_		 190 190						aanı			
SEQ	Ö		646		646			646						646			V V	3		+	040					245				-

PDB annotation	SPECIFICITY																													
Coumpound		IMMUNOGLOBULIN 3D6 FAB IDFB 3	IMMUNOGLOBULIN FV FRAGMENT OF A	HUMANIZED VERSION OF	THE ANTI-CD18 1FGV 3	AN IIBOD I H32' (HUH32- AA FV) 1FGV 4	IMMUNOGLOBULIN FV	FRAGMENT OF A	HUMANIZED VERSION OF THE ANTI-CD18 1FGV 3	ANTIBODY 'H52' (HI JH52-	AA FV) IFGV 4	IMMUNOGLOBULIN FV	FRAGMENT OF	HUMANIZED ANTIBODY	4D5, VERSION 8 IFVC 3	IMMUNOGLOBULIN FV	FRAGMENT OF	HUMANIZED ANTIBODY	4D5, VERSION 8 1FVC 3	IMMUNOGLOBULIN FAB	FRAGMENT OF	HUMANIZED ANTIBODY	4D5, VERSION 4 1FVD 3	COMPLEX(ANTIBODY-	ANTIGEN) FV FRAGMENT	(IGG1, KAPPA) (LIGHT	AND HEAVY VARIABLE	DOMAINS 1JHL 3 NON-	COVALENTLY	ASSOCIATED) OF
SeqFold Score							57.39			_						53.42								52.05						
PMF Score		1.00	0.94		-							86.0						-		00.1							•			
Verify Score		0.22	0.43									0.33								0.32										
PSI BLAST Score		6.8e-50	3.4e-53				3.4e-53					3.4e-50				3.4e-50			_	1.2e-50				1.5e-44						
End		126	126				129					126				130				126	-			130						
Start AA		20	70				20					20				20				20				20						
Chain ID		L	7				7					٧				٧				4				T						
PDB ID		1dfb	lfgv				lfgv					1fvc				1fvc				Ifvd		_		IjhI						
SEQ ID NO:		646	646				646					646				646				949				646	_					

PDB annotation		COMPLEX (HYDROLASE/IMMUNOGLOBULIN)	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL			COMPLEX (MHCVVRAL PEPTIDE/RECEPTOR) HLA-A2 HEAVY CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR
Coumpound	MONOCLONAL ANTI-HEN EGG 11HL 4 LYSOZYME ANTIBODY D11.15 COMPLEX WITH PHEASANT EGG 11HL 5 LYSOZYME 11HL 6	N9 NEURAMINIDASE; INMB 4 CHAIN: N; INMB 5 FAB NC10; INMB 9 CHAIN; L, H; INMB 10	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	IMMUNOGLOBULIN WAT, A VARIABLE DOMAIN FROM IMMUNOGLOBULIN LIGHT-CHAIN 1WTL 3 (BENCE-JONES PROTEIN) 1WTL 4	IMMUNOGLOBULIN FAB FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN:
SeqFold Score		52.88	58.98	50.85		119.10
PMF Score					1.00	
Verify Score					0.28	
PSI BLAST Score		5.1e-45	5.1e-40	3.4e-49	1e-53	1.7e-40
End		130	130	129	126	135
Start AA		20	21	20	20	24
Chain ID		1	А	Ą	7	Ω
PDB UD		Inmb	Itcr	l wtl	2fgw	1ao7
SEQ B B		646	646	646	646	648

1807 D 25 137 1.76-40 0.45 1.00 H.A-A 0201; CHAIN: 4; BETA-2 MICROGLOBULIN; CHAIN: BT TAX PEPTIDE; CHAIN: BT TAX PEPTIDE; CHAIN: CT T CELL RECEPTOR ALPHA; CHAIN: CT T CELL RECEPTOR BETA; CHAIN: BT T CELL RECEPTOR BETA; CHAIN: BT T CELL RECEPTOR BETA; CHAIN: BT T CELL RECEPTOR V-AB; CHAIN: CT T CELL RECEPTOR V-AB; CHAIN: CT CELL RECEPTOR ALPHA; CHAIN: CT CELL RECEPTOR ALPHA; CHAIN: CT CELL RECEPTOR ALPHA; CHAIN: CT CELL RECEPTOR ALPHA; CHAIN: CT CELL RECEPTOR ALPHA; CHAIN: CHAIN: CT CELL RECEPTOR ALPHA; CHAIN: CHAIN: CT CELL RECEPTOR BETA; CHAIN: CT CELL RECEPTOR BETA; CHAIN: CHAIN: CHAIN: CHAIN: CHAIN: A, CHAIN: CT CELL RECEPTOR DIO (ALPHA-BETA T CELL RECEPTOR DIO (ALPHA-BETA T CELL RECEPTOR DIO (ALPHA-CHAIN); CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: CHAIN: A, CHAIN: B, CHAIN: CHAIN: CHAIN: A, CHAIN: B, CHAIN: CHAIN: CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: B, CHAIN: CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: A, CHAIN: B, CHAIN: CHAIN: A, CHAIN: A, CHAIN: B, CHAIN: CHAIN: A, CHAIN: A, CHAIN: B, CHAIN: CHAIN: A, CHAIN: B, CHAIN: CHAIN: A, CHAIN: B, CHAIN: B, CHAIN: CHAIN: A, CHAIN: B, CHAIN:	SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
1807 D 25 137 1.76-40 0.45 1.00 H.A-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PETTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: B; T CELL RECEPTOR ALPHA; CHAIN: B; T CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: B; T CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: A; B; CHAIN: B; TAX PETTIDE; CHAIN: B; TAX PETTIDE; CHAIN: B; TAX PETTIDE; CHAIN: B; TAX PETTIDE; CHAIN: B; T CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: CHAIN: C, T CELL RECEPTOR ALPHA; CHAIN: C, T CELL RECEPTOR ALPHA; CHAIN: C, T CELL RECEPTOR BETA; CHAIN: B; T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; CHAIN: C, T CELL RECEPTOR DIUR; C, T CELL RECEPTOR DIURC; C, T CELL RECEPTOR DIURC; C, T CELL RECEPTOR DIURC; C, T CELL RECEPTOR DIURC; C, T C, T C, T C, T C, T C, T C, T C,										斑	
164 154 1642 17.78 T.CELL RECEPTOR ALPHA; 1642 131 1.76-42 17.78 T.CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: B. T.CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. A. B. B. A. B. B. B. A. B. B. B. B. A. B. B. B. B. B. B. B. B. B. B. B. B. B.	∞	lao7	Ω	25	137	·	0.45	1.00		HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B: TAX PEPTIDE:	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA-A2 HEAVY CHAIN: CI ASS I MHC. T-CELL
1688 A 23 131 1.7e-42 77.78 TCELL RECEPTOR BETA; CHAIN: Biss A 24 133 1.7e-42 0.48 1.00 TCELL RECEPTOR V-ALPHA DOMAIN; CHAIN: A B; A B; BETA-2 MICROGLOBULIN; CHAIN: B TAX PEPTIDE; CHAIN: B TAX PEPTIDE; CHAIN: B TAX PEPTIDE; CHAIN: B TAX PEPTIDE; CHAIN: B TAX PEPTIDE; CHAIN: CHAIN: B TCELL RECEPTOR ALPHA; CHAIN: B TCELL RECEPTOR BETA; CHAIN: B TCELL RECEPTOR DIO; CHAIN: CHAIN										CHAIN: C; T CELL RECEPTOR AI PHA:	RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL)
A 23 131 1.7e-42 77.78 T CELL RECEPTOR V- ALPHA DOMAIN; CHAIN: A, B; D 24 137 1e-42 0.53 1.00 T CELL RECEPTOR V- ALPHA DOMAIN; CHAIN: A, B; CHAIN: C, T CELL RECEPTOR V- ALPHA DOMAIN; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: C, T CELL RECEPTOR ALPHA; CHAIN: D, T CELL RECEPTOR ALPHA; CHAIN: D, T CELL RECEPTOR BETA; CHAIN: E, CHAIN: D, T CELL RECEPTOR (TCR) (D10); CHAIN: A, B; A 25 134 3.4e-43 0.34 1.00 ALPHA-BETA T CELL RECEPTOR (TCR) (D10); CHAIN: A, B; A 25 134 3.4e-43 0.34 1.00 ALPHA CHAIN; CHAIN: A, B; T-CELL RECEPTOR D10 CHAIN: B, E, MHC L-AK A						,				CHAIN: D; T CELL	PEPTIDE/RECEPTOR
1688 A 23 131 1.76-42 77.78 T CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: A, B; A 133 1.76-42 0.48 1.00 T CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: A, B; A 137 16-42 0.53 1.00 HLA-A 0201; CHAIN: A; B ET A-2 MICROGLOBULIN; CHAIN: B; T AY PEPTIDE; CHAIN: B; T CELL RECEPTOR ALPHA; CHAIN: B; T CELL RECEPTOR D10; CHAIN: B; T CELL RECEPTOR D10; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: CHAIN: A; CHAIN: CHAIN: A; CHAIN: CHAIN: CHAIN: CHAIN: A, B; T CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN: A, B; T CELL RECEPTOR D10 (BETA CHAIN); CHAIN: B, E, MHC L-AK A										RECEPTOR BETA; CHAIN: E;	
1688 A 24 133 1.76-42 0.48 1.00 TCELL RECEPTOR V. ALPHA DOMAIN; CHAIN: A, B; B; A 137 16-42 0.53 1.00 HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B, TAX PEPTIDE; CHAIN: B, TAX PEPTIDE; CHAIN: B, TAX PEPTIDE; CHAIN: B, TAX PEPTIDE; CHAIN: B, TAX PEPTIDE; CHAIN: B, TAX PEPTIDE; CHAIN: B, TCELL RECEPTOR BETA; CHAIN: B, CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: CHAIN: CHAIN: A, E, T-CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN: B, E, MHC L-AK A, CHAIN:	<u>&</u>	1688	A	23	131	1.7e-42			77.78	T CELL RECEPTOR V.	T CELL RECEPTOR TCR; T CELL RECEPTOR, MHC CLASS I, HUMAN
1b88 A 24 133 1.7e-42 0.48 1.00 TCELL RECEPTOR V-ALPHA DOMAIN; CHAIN: A, B; A, B; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: B; TAX PEPTIDE; CHAIN: CHAIN: D; T CELL RECEPTOR (TCR) (D10); CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN: B, F, MHC L-AK A		-								A, B;	IMMUNODEFICIENCY VIRUS, 2 MOLECULAR RECOGNITION
1bd2 D 24 137 1e-42 0.53 1.00 HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: B; TAX PEPTIDE; CHAIN: A; B; T-CELL RECEPTOR D10 (ALPHA-BETA T CELL RECEPTOR D10; CHAIN: A; E; T-CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN: A, E; T-CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN: B, F; MHC L-AK A	œ,	1688	4	24	133	1.7e-42	0.48	1.00		T CELL RECEPTOR V.	T CELL RECEPTOR TCR; T CELL RECEPTOR MHC CLASS I HIMAN
1bd2 D 24 137 1e-42 0.53 1.00 HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: B; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E; CHAIN: B; CHAIN: C; CAIN: CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: A; CHAIN: CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; CHAIN; B, F; MHC L-AK A										A, B;	IMMUNODEFICIENCY VIRUS, 2 MOLECULAR RECOGNITION
16 14 15 15 15 15 16 17 17 18 17 18 17 18 17 18 17 18 17 18 17 18 18	∞	1bd2	Ω	24	137	1e-42	0.53	1.00		HLA-A 0201; CHAIN: A;	COMPLEX (MHC/VIRAL
Ibwm A 24 134 3.4e-44 0.22 1.00 ALPHA-BETA T CELL RECEPTOR BETA; CHAIN: B; CHAIN: B; CHAIN: B; CHAIN: A; CHAIN: A; CHAIN: A, B; T-CELL RECEPTOR (TCR) (D10); CHAIN: A, B; T-CELL RECEPTOR D10 ALPHA-CHAIN; CHAIN: A, B; T-CELL RECEPTOR D10 CHAIN: B, F; MHC I-AK A CHAIN: CHAIN: B, F; MHC I-AK A CHAIN: CHAIN: B, F; MHC I-AK A CHAIN: CHAIN: B, F; MHC I-AK A CHAIN:								_		BELA-2 MICKUGLOBULIN; CHAIN: B: TAX PEPTIDE:	FEFTIDE/RECEFTOR) HEA AZ HEAV I CHAIN: COMPLEX (MHC/VIRAL
December 1984 December 1984 December 1984 December 1984 December 1984 December 1984 December 1984 December 1984 December 1984 December 1985 December 1985 December 1986 December 198						•				CHAIN: C; T CELL	PEPTIDE/RECEPTOR)
1bwm A 24 134 3.4e-44 0.22 1.00 ALPHA-BETA T CELL						,				RECEPTOR ALPHA; CHAIN: D: T CFLI	
1bwm A 24 134 3.4e-44 0.22 1.00 ALPHA-BETA T CELL RECEPTOR (TCR) (D10); CHAIN: A; CHAIN: A; CHAIN: B, E. T-CELL RECEPTOR D10 CHAIN: B, E. MHC I-AK A CHAIN: B, F. MHC I-AK A CHAIN: B,									,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	RECEPTOR BETA; CHAIN: E:	
RECEPTOR (TCR) (D10); CHAIN: A; CHAIN: A; CHAIN: A; CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN: A, E; T-CELL RECEPTOR D10 (BETA CHAIN); CHAIN: B, F; MHC I-AK A	∞	1bwm	¥	24	134	3.4e-44	0.22	1.00		ALPHA-BETA T CELL	IMMUNE SYSTEM
1d9k A 25 134 3.4e-43 0.34 1.00 T-CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN); CHAIN: A, E; T-CELL RECEPTOR D10 (BETA CHAIN); CHAIN: B, F; MHC I-AK A						,				RECEPTOR (TCR) (D10); CHAIN: A;	IMMUNORECEPTOR, IMMUNE
(ALPHA CHAIN); CHAIN: A, E; T-CELL RECEPTOR DI0 (BETA CHAIN); CHAIN: B, F; MHC I-AK A		1907	4	3,5	122	3 10 13	0.34	5		T.CELL PECEPTOR DIO	SYSIEM NAME LAK: MHC I.
D10 (BETA CHAIN); CHAIN: B, F; MHC I-AK A		4	τ	3	<u> </u>	C+-3+'C	<u> </u>	2		(ALPHA CHAIN); CHAIN: A, E; T-CELL RECEPTOR	AK; T-CELL RECEPTOR, MHC CLASS II, D10, I-AK
										D10 (BETA CHAIN); CHAIN: B, F; MHC I-AK A	

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
NO:	a į		AA	AA	BLAS1 Score	Score	Score	Score		
									CHAIN (ALPHA CHAIN); CHAIN: C, G; MHC I-AK B	
									CHAIN (BETA CHAIN); CHAIN: D, H;	
									CONALBUMIN PEPTIDE; CHAIN: P, O;	
648	1kb5	A	24	133	1e-44	89.0	1.00		KB5-C20 T-CELL ANTIGEN	COMPLEX
									RECEPTOR; CHAIN: A, B;	(IMMUNOGLOBULIN/RECEPTOR) TCR
					_				ANTIBODY DESIRE-1; CHAIN: L, H;	VAPLHA VBETA DOMAIN; I-CELL RECEPTOR, STRAND SWITCH, FAB,
										ANTICLONOTYPIC, 2 (IMMINOGLOBULIN/RECEPTOR)
648	1kb5	A	24	135	le-44			74.20	KB5-C20 T-CELL ANTIGEN	COMPLEX
					112				RECEPTOR; CHAIN: A, B; ANTIRODY DESIRE-1:	(IMMUNOGLOBULIN/RECEPTOR) TCR
									CHAIN: L, H;	RECEPTOR, STRAND SWITCH, FAB,
										ANTICLONOTYPIC, 2 (IMMI INOG! OBLIT IN/RECEPTOR)
648	Ig1	Ω	24	142	1.7e-40			74.24	MHC CLASS I HLA-A;	IMMUNE SYSTEM HUMAN
									CHAIN: A; BETA-2	TCR/PEPTIDE/MHC COMPLEX, HLA-
									MICROGLOBULIN; CHAIN:	A2, HTLV-1, TAX, TCR, T 2 CELL
									B; I AX PEP IIDE P6A;	KECEPIUK, IMMUNE SYSIEM
						· . <u>-</u>			RECEPTOR: CHAIN: D:	
			İ						HLA-A 0201; CHAIN: E;	
648	lqrn	Ω	25	137	1.7e-40	0.49	1.00		MHC CLASS I HLA-A;	IMMUNE SYSTEM HUMAN
						_			CHAIN: A; BETA-2	TCR/PEPTIDE/MHC COMPLEX, HLA-
	-					_			MICROGLOBULIN; CHAIN:	A2, HTLV-1, TAX, TCR, T 2 CELL
									B; TAX PEPTIDE P6A;	KECEPTOR, IMMONE SYSTEM
									CHAIN: C, FIMAIN 1-CELL	•
							_		KECEPTOK; CHAIN: D; HLA-A 0201; CHAIN: E;	
649	1a4j	L	21	238	le-73			82.63	IMMUNOGLOBULIN,	IMMUNOGLOBULIN

SEQ ID NO:	PDB ID	Chain ID	Start AA	End AA	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									DIELS ALDER CATALYTIC ANTIBODY; CHAIN: L, H, A, B;	IMMUNOGLOBULIN, ANTIBODY, CATALYTIC ANTIBODY, DIELS ALDER, 2 GERMLINE
649	lae6	J	21	229	8.5e-73			83.03	ANTIBODY CTM01; CHAIN: L, H;	IMMUNOGLOBULIN, FAB IMMUNOGLOBULIN, FAB FRAGMENT, HUMANISATION
649	1b2w	1	20	229	5.1e-85	0.01	0.58		ANTIBODY (LIGHT CHAIN); CHAIN: L; ANTIBODY (HEAVY CHAIN); CHAIN: H;	IMMUNE SYSTEM IMMUNOGLOBULIN; IMMUNOGLOBULIN; ENGINEERING, HUMANIZED AND CHIMERIC ANTBODY, FAB, 2 X-RAY STRUCTURE, THREE-DIMENSIONAL STRYCTURE, GAMMA- 3
649	1b2w	7	21	240	5.1e-85			85.20	ANTIBODY (LIGHT CHAIN); CHAIN: L; ANTIBODY (HEAVY CHAIN); CHAIN: H;	INTERFERON, IMMUNE SYSTEM IMMUNE SYSTEM IMMUNOGLOBULIN; IMMUNOGLOBULIN; ENGINEERING, HUMANIZED AND CHIMERIC ANTIBODY, FAB, 2 X-RAY STRUCTURE, THREE-DIMENSIONAL STRYCTURE, GAMMA- 3 NYTERFERON IMMINE SYSTEM
649	164j	J	21	240	5.1e-79			81.22	ANTIBODY; CHAIN: L, H;	ANTIBODY ENGINEERING ANTIBODY ENGINEERING HUMANIZED AND CHIMERIC ANTIBODIES, 2 FAB, X-RAY STRUCTURES, GAMMA-INTERFERON
649	1b6d	A	70	227	5.1e-84	0.15	0.70		IMMUNOGLOBULIN; CHAIN: A, B;	IMMUNOGLOBULIN IMMUNOGLOBULIN, KAPPA LIGHT- CHAIN DIMER HEADER
649	16j1	1	20	228	3.4e-86	-0.04	0.54		FAB FRAGMENT; CHAIN: L, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR; CHAIN: V, W;	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR

21 237 3.4e-82 81.96 CAMPATH-IH-LIGHT CAMPATH-HEIBEAVY CHAIN; CHAIN; CHAIN; H; PEPTIDE ANTIGEN; CHAIN: P; CAMPATH-HEIBEAVY CHAIN; CHAIN: H; PEPTIDE ANTIGEN; CHAIN: P; CHAIN: P; CHAIN: P; CHAIN: H; PEPTIDE ANTIGEN; CHAIN: H; PEPTIDE ANTIGEN; CHAIN: H; CHAIN: CHAIN: H; CHAIN: L, H; CHAIN: L, H; SHORT CHAIN; CHAIN: A, C, CHAIN: L, H; CHAIN: CHAIN: CHAIN: B, D CHAIN: L, H; CHAIN: CHAIN: CHAIN: A, C, C, C, C, C, C, C, C, C, C, C, C, C,	Chain ID		Start	End	PSI BLAST	Verify Score	PMF	SeqFold	Coumpound	PDB annotation
237 3.46-82 81.96 CAMPATH-IH:LIGHT CHAIN; CHAIN: L; CAMPATH-IH:HEAVY CHAIN; CHAIN: H; PEPTIDE ANTIGEN; CHAIN: P; CHAIN: CHAIN: CHAIN: A, C, C, C, T C F A B FRAGMENT; CHAIN: CHAIN: CHAIN: B, D CHAIN: P; CHAIN: CHAIN: CHAIN: A, C, C, C, T C F A B FRAGMENT; CHAIN: CHAIN: CHAIN: B, D CHAIN: P; CHAIN: CHAIN: CHAIN: B, D CHAIN: CHAIN: CHAIN: B, D CHAIN: CH					Score			3		
229 3.4e-75 82.55 IGG FAB (IGG3, KAPPA); CHAIN: H; PRETIDIE ANTIGEN; CHAIN: H; PRETIDIE ANTIGEN; CHAIN: P; CHAIN: P; CHAIN: P; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: CHAIN	L 21	21		237	3.4e-82			81.96	CAMPATH-1H:LIGHT CHAIN; CHAIN: L;	ANTIBODY THERAPEUTIC, ANTIBODY, CD52
229 3.4e-75 82.55 IGG FAB (HUMAN IGGI, KAPPA); CHAIN: L, H; KAPPA); CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: B, D IGM RF 2A2; CHAIN: CHAIN: CHAIN: CHAIN: L, H; IGM RF 2A2;									CAMPATH-IH:HEAVY CHAIN: CHAIN: H:	
229 3.4e-75 82.55 IGG FAB (HUMAN IGGI, KAPPA); CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: CHAIN: CHAIN: CHAIN: A, C; 7C8 FAB FRAGMENT; CHAIN: B, D IGM RF 2A2; CHAIN: B, D IGM RF 2A2; CHAIN: B, D IGM RF 2A2; CHAIN: B, D IGM RF 2A2; CHAIN: B, CHAIN:									PEPTIDE ANTIGEN; CHAIN: P;	
229 3.4e-75 82.55 IGG FAB (IGG3, KAPPA); CHAIN: L, H; 229 1.7e-74 81.49 7C8 FAB FRAGMENT; 229 3.4e-88 -0.04 0.84 IGM RF 2A2; CHAIN: B, D 1.5e-80 I.2e-80 B1.24 IMMUNOGLOBULIN FAB 529 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; H; 229 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; Himmunoglobulin NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; IMMUNOGLOBULIN NMC-4 IGG1; IMMUNOGLOBULIN NMC-4 IG	L 22	22		240	5.1e-73			84.78	IGG FAB (HUMAN IGGI,	IMMUNOGLOBULIN CBR96 FAB
229 3.46-75 82.55 IGG FAB (IGG3, KAPPA); CHAIN: L, H; CHAIN: L, H; SHORT CHAIN; CHAIN: A, C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C; 7C8 FAB FRAGMENT; CHAIN: B, D C; 7C8 FAB FRAGMENT; CHAIN: B, D C; 7C8 FAB FRAGMENT; CHAIN: B, D C; 7C8 FAB FRAGMENT; CHAIN: B, D C; 7C8 FAB FRAGMENT; CHAIN: CHAIN: L, H; CHAIN: L, H; CHAIN: L, H; CHAIN: L; IMMUNOGLOBULIN NMC-A GO CHAIN: L; IMMUNOGLOBULIN		=							KAPPA); CHAIN: L, H;	(IMMUNOGLOBULIN); IMMUNOGLOBULIN.
229 3.4e-75 82.55 IGG FAB (IGG3, KAPPA); CHAIN: L, H; CHAIN: L, H; SHORT CHAIN; CHAIN: A, C; 7C8 FAB FRAGMENT; LONG CHAIN: CHAIN: B, D C; 7C8 FAB FRAGMENT; LONG CHAIN: A, C, E; IGM RF 2A2; CHAIN: B, D C; 7C8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAIN: CGAIN: CGAIN: C, TC8 FAB FRAGMENT; CGAIN: CGAI										IMMUNOGLOBULIN C REGION, GLYCOPROTEIN, ANTIB
229 1.7e-74 81.49 7C8 FAB FRAGMENT; SHORT CHAIN; CHAIN: A, C, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D F; IMMUNOGLOBULIN G, B, CHAIN: G, H; CHAIN: G, H; CHAIN: C, H; CHAIN: C, H; CHAIN: C, H; CHAIN: L, H; CASP 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC-4 1GG1; CHAIN: L; IMMUNOGLOBULIN N	T 20	70		229	3.4e-75			82.55	IGG FAB (IGG3, KAPPA);	IMMUNOGLOBULIN MBR96 FAB
229 1.7e-74 81.49 7C8 FAB FRAGMENT; SHORT CHAIN; CHAIN: A, C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D LONG CHAIN; CHAIN: B, D IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: L, H; E; IGG1; CHAIN: L, H; E; IGG1; CHAIN: L, L, L, L, L, L, L, L, L, L, L, L, L,									CHAIN: L, H;	(IMMUNOGLOBULIN);
229 1.7e-74 81.49 7C8 FAB FRAGMENT; SHORT CHAIN; CHAIN: A, C, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D C, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D G, 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H; CHAIN: G, H; CHAIN: G, H; SG9; CHAIN: L, H; IMMUNOGLOBULIN FAB 5G9; CHAIN: L, H; IMMUNOGLOBULIN NMC-4 1GG1; CHAIN: L; IMMUNOGLOBULIN NM						***				IMMUNOGLOBULIN C REGION, GLYCOPROTEIN, TRANSMEMBRANE
229 3.4e-88 -0.04 0.84 C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D IGM RF 2A2; CHAIN: B, C E; IGM RF 2A2; CHAIN: B, D F; IMMUNOGLOBULIN G BIDDING PROTEIN A; CHAIN: G, H; CHAIN: G, H; CHAIN: G, H; CHAIN: G, H; CHAIN: G, H; CHAIN: C, CHAIN: C,	A 2	7		229	1.7e-74			81.49	7C8 FAB FRAGMENT;	IMMUNE SYSTEM ABZYME
C; 7C8 FAB FRAGMENT; LONG CHAIN; CHAIN: B, D LONG CHAIN; CHAIN: B, D IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B, D, F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H; CHAIN: G, H; SG9; CHAIN: L, H; IMMUNOGLOBULIN NMC- A IGG1; CHAIN: L; IMMUNOGLOBULIN NM								•	SHORT CHAIN; CHAIN: A,	TRANSITION STATE ANALOG,
229 3.4e-88 -0.04 0.84 IGM RF 2A2; CHAIN; D. C									C; 7C8 FAB FRAGMENT;	IMMUNE SYSTEM
229 3.4e-88 -0.04 0.84 IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B, D, F; IMMUNOGLOBULIN G BINDING PROTEIN A; CHAIN: G, H; CHAIN: G, H; SG9; CHAIN: L, H; SG9; CHAIN: L, H; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; VIONI	1	- 1'							LONG CHAIN; CHAIN: B, D	
229 1.2e-80 81.24 IMMUNOGLOBULIN FAB 5G9; CHAIN: I., H; MMUNOGLOBULIN FAB 5G9; CHAIN: I., H; MMUNOGLOBULIN NMC-4 1GG1; CHAIN: I., IIMMUNOGLOBULIN NMC-4 1GG1; CHAIN: I., NON-4 1GG1; CHAIN:	A	7	0	229	3.4e-88	-0.04	0.84		IGM RF 2A2; CHAIN: A, C, E; IGM RF 2A2; CHAIN: B,	IMMUNE SYSTEM FAB-IBP COMPLEX CRYSTAL STRUCTURE 2.7A
229 1.2e-80 81.24 IMMUNOGLOBULIN FAB 5G9; CHAIN: L, H; MMUNOGLOBULIN NMC-229 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; VIONI									D, F; IMMUNOGLOBULIN	RESOLUTION BINDING 2 OUTSIDE
229 1.2e-80 81.24 IMMUNOGLOBULIN FAB 5G9; CHAIN: L, H; 5G9; CHAIN: L, H; 124 IMMUNOGLOBULIN NMC-127 0.75 IMMUNOGLOBULIN NMC-14 1GG1; CHAIN: L; IMMUNOGLOBULIN NMC-14 1GG1; CHAIN: L; IMMUNOGLOBULIN NMC-14 1GG1; CHAIN: L; IMMUNOGLOBULIN NMC-14 1GG1; CHAIN: L; VON									G BINDING PROTEIN A;	THE ANTIGEN COMBINING SITE
229 1.2e-80 81.24 IMMUNOGLOBULIN FAB 5G9; CHAIN: L, H; 5G9; CHAIN: L, H; 129 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC-16G1; CHAIN: L; IMMUNOGLOBULIN NMC-16G1; CHAIN: L; IMMUNOGLOBULIN NMC-16G1; CHAIN: L; IMMUNOGLOBULIN NMC-16G1; CHAIN: L; NON-16G1;									CHAIN: G, H;	SUPERANTIGEN FAB VH3 3 SPECIFICITY
229 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-4 IGG1; CHAIN: L; VION	7		12	229	1.2e-80			81.24	IMMUNOGLOBULIN FAB	IMMUNOGLOBULIN FAB, FAB LIGHT
229 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC- 4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-					_				5G9; CHAIN: L, H;	CHAIN, FAB HEAVY CHAIN;
229 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC- 4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-										ANTIBODY, FAB, ANTI-TF,
229 3.4e-84 -0.27 0.75 IMMUNOGLOBULIN NMC- 4 IGG1; CHAIN: L; IMMUNOGLOBULIN NMC-										MONOCLONAL, MURINE, IMMIJNOGI, OBIJI, IN
	L 2	12	9	229	3.4e-84	-0.27	0.75		IMMUNOGLOBULIN NMC-	IMMUNE SYSTEM VON WILLEBRAND
									4 IGG1; CHAIN: L;	FACTOR, GLYCOPROTEIN IBA
								•••	MMCNOGLOBULIN NMC-	(A:ALPHA) BINDING, 2 COMPLEX (WIT I PER AND/IMMINOGIORIII IN)

Chain S	S	Start	End	PSI BI AST	Verify	PMF	SeqFold	Coumpound	PDB annotation
Y.	ri .	•	5	Score	Score	arore	Score		
								WILLEBRAND FACTOR; CHAIN: A;	BLOOD COAGULATION TYPE 3 2B VON WILLEBRAND DISEASE
A 20 229		22	0,	5.1e-85	0.27	0.75	-	IMMUNOGLOBULIN FAB FRAGMENT OF HUMANIZED ANTIBODY 4D5. VERSION 4 1FVD 3	
L 21 237		23	7	6.8e-80			82.59	ENVELOPE PROTEIN GP120; CHAIN: G; CD4; CHAIN: C; ANTIBODY 17B; CHAIN: L, H;	COMPLEX (HIV ENVELOPE PROTEIN/CD4/FAB) COMPLEX (HIV ENVELOPE PROTEIN/CD4/FAB), HIV-1 EXTERIOR 2 ENVELOPE GP120, T- CELL SURFACE GLYCOPROTEIN CD4, 3 ANTIGEN-BINDING FRAGMENT OF HUMAN IMMUNOGLOBULIN 17B, 4 GLYCOSYLATED PROTEIN
A 20 229		22	_	1e-84	-0.01	0.53		IGG2A INTACT ANTIBODY - MAB231; CHAIN: A, B, C, D	IMMUNOGLOBULIN INTACT IMMUNOGLOBULIN V REGION C REGION, IMMUNOGLOBULIN
A 21 229		225		1.5e-66			81.70	LAMBDA III BENCE JONES PROTEIN CLE; CHAIN: A, B	IMMUNÓGLOBULIN IMMUNOGLOBULIN, BENCE JONES PROTEIN
D 21 226		226		1.7e-55			224.26	MHC CLASS I HLA-A; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE P6A; CHAIN: C; HMAN T-CELL RECEPTOR; CHAIN: D; HLA-A 0201; CHAIN: E:	IMMUNE SYSTEM HUMAN TCR/PEPTIDE/MHC COMPLEX, HLA- A2, HTLV-1, TAX, TCR, T 2 CELL RECEPTOR, IMMUNE SYSTEM
L 20 229		22	6	8.5e-84	-0.03	0.65		MONOCLONAL ANTIBODY 3A2; CHAIN: H, L;	MONOCLONAL ANTIBODY MONOCLONAL ANTIBODY, FAB- FRAGMENT, REPRODUCTION
A 21 231	,,	23		2.7e-74			237.84	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL
L 22 229	П	22		8.5e-84	0.14	0.62		TR1.9 FAB; CHAIN: L, H;	IMMUNOGLOBULIN TR1.9, ANTI-

PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									THYROID PEROXIDASE, AUTOANTIBODY, 2 IMMUNOGLOBULIN
		70	229	3.4e-87	-0.04	0.52		IMMUNOGLOBULIN FAB	
								HUMANIZED VERSION OF	
								THE ANTI-CD18 2FGW 3	
								ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4	
	T	21	229	3.4e-87			83.47	IMMUNOGLOBULIN FAB	
								FRAGMENT OF A	
							-	HUMANIZED VERSION OF	
								ANTIRODY 'H52' (HITHS2.	
Į.								OZ FAB) 2FGW 4	
	1	21	229	3.4e-80			81.24	IMMUNOGLOBULIN	
				-			-	ANTIGEN-BINDING	
					_			FRAGMENT OF THE MIRINF ANTI-	
								PHENYLARSONATE 6FAB	
								3 ANTIBODY 36-71, FAB 36-71 6FAB 4	
Ιl									
lae6	Ħ	21	245	1.4e-91	0.55	1.00		ANTIBODY CTM01;	IMMUNOGLOBULIN
								CHAIN: L, H;	IMMUNOGLOBULIN, FAB FRAGMENT, HUMANISATION
	н	21	247	5.1e-93	0.57	00.1		HUMAN	COMPLEX (VIRAL
								IMMUNODEFICIENCY	CAPSID/IMMUNOGLOBULIN) HIV-1
				-				VIRUS TYPE 1 CAPSID	CA, HIV CA, HIV P24, P24; FAB, FAB
								CHAIN: A, B; ANTIBODY	LIGHT CHAIN, FAB HEAVY CHAIN
								FAB25.3 FRAGMENT;	COMPLEX (VIRAL
								CHAIN: H, K, L, M;	CAPSID/IMMUNOGLOBULIN), HIV,
1	Э	23	264	8.5e-62			303.44	HLA-A 0201; CHAIN: A;	COMPLEX (MHC/VIRAL

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
9 ö	9	8	AA	¥	BLAST Score	Score	Score	Score		
									BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C: T CELL	PEPTIDE/RECEPTOR) HLA-A2 HEAVY CHAIN; CLASS I MHC, T-CELL RECEPTOR VIRAI PEPTIDE 2
									RECEPTOR ALPHA;	COMPLEX (MHC/VIRAL
									CHAIN: D; T CELL	PEPTIDE/RECEPTOR
									RECEPTOR BETA; CHAIN: F:	
650	1bd2	E	23	264	1.7e-84			397.14	HLA-A 0201; CHAIN: A;	COMPLEX (MHC/VIRAL
									BETA-2 MICROGLOBULIN;	PEPTIDE/RECEPTOR) HLA A2 HEAVY
									CHAIN: B; TAX PEPTIDE;	CHAIN; COMPLEX (MHC/VIRAL
								_	CHAIN: C; T CELL	PEPTIDE/RECEPTOR)
									RECEPTOR ALPHA;	
									CHAIN: D; T CELL	
									RECEPTOR BETA; CHAIN:	
									E;	
650	1bec		23	264	5.4e-95			324.99	14.3.D T CELL ANTIGEN	RECEPTOR T CELL RECEPTOR 1BEC
									RECEPTOR; 1BEC 5	14
									CHAIN: NULL; 1BEC 6	
650	1bec		24	263	5.4e-95	0.73	1.00		14.3.D T CELL ANTIGEN	RECEPTOR T CELL RECEPTOR 1BEC
									RECEPTOR; 1BEC 5	
									CHAIN: NULL; 1BEC 6	
650	1dzb	٧	6	134	8.5e-35	0.34	0.82		SCFV FRAGMENT 1F9;	COMPLEX (ANTIBODY ANTIGEN) 1,4-
									CHAIN: A, B; TURKEY	BETA-N-ACETYLMURAMIDASE C;
_									EGG-WHITE LYSOZYME	SINGLE-DOMAIN ANTIBODY,
									C; CHAIN: X, Y;	TURKEY EGG-WHITE LYSOZYME, 2
							•			ANTIBODY-PROTEIN COMPLEX,
										SINGLE-CHAIN FV FRAGMENT
920	1e60	н	21	247	5.1e-92	0.49	1.00		IMMUNOGLOBULIN	IMMUNOGLOBULIN FAB, ANTIBODY,
									LIGHT CHAIN; CHAIN: L;	ANTIGEN, HIV-1, P24, CA
									IMMUNOGLOBULIN	
									HEAVY CHAIN; CHAIN: H;	
650	<u>1</u>	æ	17	134	1.5e-33	0.11	0.89		ACETYLCHOLINE	IMMUNE SYSTEM IG-FOLD, IMMUNO
									RECEPTOR ALPHA;	COMPLEX, ANTIBODY-ANTIGEN,
									CHAIN: A; FV ANTIBODY	BEIA-IUKN

		_	_			_		_		_					_						_	_	_	_	_			_						
PDB annotation																							IMMUNOGLOBULIN INTACT	IMMUNOGLOBULIN V REGION C									IMMUNOGLOBULIN	IMMUNOGLOBULIN,
Coumpound		FRAGMENT; CHAIN: B;	COMPLEX	(ANTIBODY/ANTIGEN)	FAB FRAGMENT OF THE	MONOCLONAL	ANTIBODY F9.13.7 (IGG1)	1FBI 3 COMPLEXED WITH	LYSOZYME (E.C.3.2.1.17)	1FBI 4	IMMUNOGLOBULIN FAB	FRAGMENT OF	HUMANIZED ANTIBODY	4D5, VERSION 4 IFVD 3	COMPLEX	(ANTIBODY/BINDING	PROTEIN) IGG1 FAB	FRAGMENT COMPLEXED	WITH PROTEIN G	(DOMAIN III) 1IGC 5	PROTEIN G,	STREPTOCOCCUS 11GC 15	IGG2A INTACT ANTIBODY	- MAB231; CHAIN: A, B, C,	IMMUNOGLOBULIN ANTI-	PHOSPHATIDYLINOSITOL	SPECIFIC	PHOSPHOLIPASE C	DIABODY 1LMK 3	SYNONYMS: L5MK16	DIABODY, SINGLE-CHAIN	FV DIMER 1LMK 4	NIG9 (IGG1=LAMBDA=);	CHAIN: L, H;
SeqFold	Score																										-	-	•					
PMF	Score		0.99								1.00				1.00								1.00		0.53								1.00	
Verify	Score		0.48			_					0.38				0.42								0.28		0.16								99.0	
PSI	BLAST Score		5.1e-91								5.1e-92		•		le-92		_				_		5.1e-95		6.8e-32								1.4e-94	
<u> </u>	AA		247								250				250								258		134								247	
	AA		21								21				21								21		∞								21	
Chain	er .		н								В				E								Ф		A								Ħ	
PDB	ar		1fbi] Lvd			:	ည္တို						_		ligt		Ilmk		_						lngp	
SEQ	NO.		029				_				059			3	050		_						029		920								650	

				_			_			
PDB annotation	IMMUNOGLOBULIN VARIABLE HEAVY (VH) DOMAIN, VARIABLE LIGHT (VL) ANTIBODY FRAGMENT, MULTIVALENT ANTIBODY, DIABODY, DOMAIN 2 SWAPPING, IMMINOGLOBILLIN	IMMUNOGLOBULIN IMMUNOGLOBULIN, SINGLE-CHAIN FV, ANTI-CARCINOEMBRYONIC 2 ANTIGEN	COMPLEX (ANTIBODY/ELECTRON TRANSPORT) FAB E8; CYT C, ANTIGEN; IMMUNOGLOBULIN, IGGI KAPPA, FAB FRAGMENT, HORSE 2 CYTOCHROME C, COMPLEX (ANTIBODY/ELECTRON TRANSPORT)					OXIDOREDUCTASE FATTY ACID HYDROXYLASE; FATTY ACID MONOOXYGENASE, HEMOPROTEIN, P450 REMARK	OXIDOREDUCTASE FATTY ACID HYDROXYLASE; FATTY ACID MONOOXYGENASE, HEMOPROTEIN, P450 REMARK	OXIDOREDUCTASE FATTY ACID HYDROXYLASE: FATTY ACID
Coumpound	SINGLE-CHAIN ANTIBODY FRAGMENT; CHAIN: A, C;	MFE-23 RECOMBINANT ANTIBODY FRAGMENT; CHAIN: A;	E8 ANTIBODY; CHAIN: L, H; CYTOCHROME C; CHAIN: F;		PHOSPHOLIPASE A2 INHIBITOR CLARA CELL 17-KDA PROTEIN ICCD 3	STEROID BINDING UTEROGLOBIN (OXIDIZED) 1UTG 4		CYTOCHROME P450; CHAIN: A, B;	CYTOCHROME P450; CHAIN: A, B;	CYTOCHROME P450; CHAIN: A, B;
SeqFold Score										114.08
PMF Score	0.92	0.98	1.00		0.43	0.09		1.00	1.00	
Verify Score	0.07	0.39	0.55		-0.41	0.07		0.40	0.40	
PSI BLAST Score	6.8e-33	8.5e-37	5.1e-91		6100.0	0.0016		2.7e-60	1.7e-45	2.7e-60
End	135	134	250		58	58		373	385	408
Start	=	=	21		1	3		_	-	
Chain ID	⋖	¥	н					A	¥	Ą
PDB ID	Inqb	lqok	l wej		lccd	lutg		1bu7	1bu7	1bu7
SEQ ID NO:	029	059	650		652	652		959	929	959

_	PDB UD	Chain ID	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
1 1										MONOOXYGENASE, HEMOPROTEIN, P450 REMARK
_	lcpt		149	390	1e-16	0.03	0.42		OXIDOREDUCTASE(OXYG ENASE) CYTOCHROME P450-TERP 1CPT 3	
) <u>~</u>	1dt6	⋖	2	388	0	0.49	1.00		CYTOCHROME P450 2C5; CHAIN: A;	OXIDOREDUCTASE PROGESTERONE 21-HYDROXYLASE, CYPIIC5 P450 1, MEMBRANE PROTEIN, PROGESTERONE 21-HYDROXYLASE, BENZO(A) 2 PYRENE HYDROXYLASE, ESTRADIOL 2-HYDROXYLASE, ESTRADIOL 2-HYDROXYLASE,
, I	1526	¥	9	373	1.3e-50	0.14	0.47		NITRIC OXIDE REDUCTASE; CHAIN: A;	OXIDOREDUCTASE NITRIC OXIDE REDUCTASE, CYTOCHROME P450NOR
∸ 1	1526	4	91	384	1.7e-07	-0.28	0.00		NITRIC OXIDE REDUCTASE; CHAIN: A;	OXIDOREDUCTASE NITRIC OXIDE REDUCTASE, CYTOCHROME P450NOR
_	1f4t	Ą	22	373	8.1e-31	-0.16	0.25		CYTOCHROME P450 119; CHAIN: A, B;	OXIDOREDUCTASE CYP119; P450 FOLD
	loxa		1	373	5.4e-61	-0.05	0.99		CYTOCHROME P450 ERYF; 10XA 5 CHAIN: NULL 10XA 6	OXIDOREDUCTASE (OXYGENASE)
	Іоха		1	399	5.4e-61			79.88	CYTOCHROME P450 ERYF; 10XA 5 CHAIN: NULL 10XA 6	OXIDOREDUCTASE (OXYGENASE)
	loxa		6	384	1.7e-19	0.26	0.95		CYTOCHROME P450 ERYF; 10XA 5 CHAIN: NULL 10XA 6	OXIDOREDUCTASE (OXYGENASE)
_ '	Iqmq	А	146	388	1.4e-06	0.03	0.27		CYTOCHROME P450; CHAIN: A;	OXIDOREDUCTASE CAMPHOR 5- MONOOXYGENASE OXIDOREDUCTASE(OXYGENASE), RU-SUBSTRATE,
1	1b34	A	_	52	1.1e-11	-0.33	0.00		SMALL NUCLEAR	RNA BINDING PROTEIN SNRNP.
									RIBONUCLEOPROTEIN SM	SPLICING, SPLICEOSOME, SM, CORE

PDB annotation		SNRNP DOMAIN, 2 SYSTEMIC LUPUS ERYTHEMATOSUS, SLE		RNA BINDING PROTEIN D3 CORE	PROTEIN SNRNP, SPLICING, SM. CORE	SNRNP DOMAIN, SYSTEMIC LUPUS 2	ERYTHEMATOSUS, SLE, RNA PRICHE DE CTEN		COMPLEX (ZINC FINGER/DNA)	FINGER, DNA-BINDING PROTEIN				COMPLEX (ZINC FINGER/DNA) COMPLEX (ZINC FINGER/DNA), ZINC	FINGER, DNA-BINDING PROTEIN			COMPLEX (ZINC FINGER/DNA) ZINC	FINGER, PROTEIN-DNA	INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX	(ZINC FINGER/DNA)	COMPLEX (ZINC FINGER/DNA) ZINC	INTERACTION, PROTEIN DESIGN, 2	CRYSTAL STRUCTURE, COMPLEX	(ZINC FINGER/DNA)
Coumpound		DI; CHAIN: A; SMALL NUCLEAR RIBONI CI FOPROTEIN SM	D2; CHAIN: B;	SMALL NUCLEAR RIBONIICI FORBOTEINI SM	D3; CHAIN: A. C. E. G. I. K:	SMALL NUCLEAR	RIBONUCLEOPROTEIN	F, H, J, L;	QGSR ZINC FINGER	DUPLEX	OLIGONUCLEOTIDE	BINDING SITE; CHAIN: B,	Ċ;	QGSR ZINC FINGER PEPTIDE; CHAIN: A;	DUPLEX	OLIGONUCLEOTIDE	BINDING SITE; CHAIN: B, C;	DNA; CHAIN: A, B, D, E;	CONSENSUS ZINC FINGER	PROTEIN; CHAIN: C, F, G;			DNA; CHAIN: A, B, D, E;	PROTEIN: CHAIN: C. F. G.		
SeqFold	Score					_						_														
PMF	Score			-0.12					0.13					0.09				0.46					0.00		-	
Verify	Score			0.14					-0.26					-0.52				-0.57					-0.52			
PSI	Score Score			5.1e-12			-		1.7e-26				_	1.7e-26				5.1e-46					1.7e-13	-		
	AA			51					244					185				244					244			
Start	AA			3					160					86				159					217			
Chain	a			В					¥					Ψ				၁		-			<u></u>			
PDB	a			143b					laih					lath				lmey					lmey			
SEQ	g ë			658					629					659				629					629			

PDB annotation	COMPLEX (TRANSCRIPTION REGULATION/DNA) COMPLEX (TRANSCRIPTION REGULATION/DNA), RNA POLYMERASE III, 2 TRANSCRIPTION INITIATION, ZINC FINGER PROTEIN	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (TRANSCRIPTION REGULATION/DNA) YING-YANG 1; TRANSCRIPTION INITIATION, INITIATOR ELEMENT, YY1, ZINC 2 FINGER PROTEIN, DNA-PROTEIN RECOGNITION, 3 COMPLEX (TRANSCRIPTION REGULATION/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA- BINDING PROTEIN/DNA)	COMPLEX (DNA-BINDING PROTEIN/DNA) FIVE-FINGER GLI; GLI, ZINC FINGER, COMPLEX (DNA-
Coumpound	TFIIIA; CHAIN: A, D; SS RIBOSOMAL RNA GENE; CHAIN: B, C, E, F;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	YYI; CHAIN: C; ADENO- ASSOCIATED VIRUS P5 INITIATOR ELEMENT DNA; CHAIN: A, B;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;	ZINC FINGER PROTEIN GLII; CHAIN: A; DNA; CHAIN: C, D;
SeqFold Score	57.65					29.77
PMF Score		0.04	0.01	0.16	0.01	
Verify Score		-0.67	-0.65	-0.44	-0.41	
PSI BLAST Score	1e-32	8.5e-30	6.8e-31	2.7e-10	6.8e-32	6.8e-32
End	273	211	244	269	273	246
Start AA	97	105	135	165	138	94
Chain ID	¥	ပ	ပ	ပ	Ą	А
PDB ID	1116	lubd	lubd	1ubd	2gli	2gli
SEQ ID NO:	629	659		659	629	659

PDB annotation	BINDING PROTEIN/DNA)		OXIDOREDUCTASE 3-ALPHA-HSD; OXIDOREDUCTASE, NAD	OXIDOREDUCTASE OXIDOREDUCTASE, ALDOSE REDUCTASE, INHIBITION, DIABETES	OXIDOREDUCTASE ALPHA/BETA TIM BARREL, PROTEIN-NADP+COMPLEX	OXIDOREDUCTASE ALDOSE REDUCTASE, INHIBITION, DIABETES	OXIDOREDUCTASE (NADP) ALDO- KETO OXIDOREDUCTASE (NADP), TIM BARREL	DE NOVO PROTEIN PROTEIN DESIGN, HYDROPHOBIC CORE, PACKING, ROTAMERS, ROC, 2 UBIQUITIN, DE NOVO PROTEIN, UBIQUITIN			UBIQUITIN UBIQUITIN, DESIGNED CORE MUTANT		IMMUNOGLOBULIN IMMUNOGLOBULIN
Coumpound		OXIDOREDUCTASE ALDOSE REDUCTASE (E.C.1.1.1.21) COMPLEX WITH NADPH 1ADS 3	3-ALPHA- HYDROXYSTEROID DEHYDROGENASE; CHAIN: A. B:	ALDOSE REDUCTASE; CHAIN: NULL;	CHO REDUCTASE; CHAIN: A;	ALDOSE REDUCTASE; CHAIN: A;	FR-1 PROTEIN; CHAIN: NULL;	ID8 UBIQUITIN; CHAIN: A;	UBIQUITIN TETRAUBIQUITIN 1TBE 3	CHROMOSOMAL PROTEIN UBIQUITIN IUBI 3	UBIQUITIN CORE MUTANT 1D7; CHAIN: A;		2E8 (IGG1=KAPPA=) ANTIBODY; CHAIN: L, H,
SeqFold Score													73.68
PMF Score		1.00	1.00	1.00	1.00	1.00	1.00	0.42	0.59	09.0	0.34		
Verify Score		0.56	0.54	0.51	0.55	0.50	0.57	-0.62	-0.48	-0.52	-0.71		
PSI BLAST Score		6.8e-42	1.7e-38	3.4e-39	3.4e-39	6.8e-42	6.8e-39	8.5e-26	le-27	1e-27	5.1e-26	•	1.7e-68
End		129	129	128	128	129	128	55	55	55	55		227
Start AA		11	∞	∞	6	11	6	-	I	1	_		18
Chain ID			A		∢	4		A	В		Ą		J
PDB ID		lads	lafs	lah4	1c9w	1el3	1frb	lc3t	1tbe	1ubi	1ud7		12e8
SEQ ID NO:		664	664	664	664	664	664	999	999	999	999		671

PDB U	Chain	Start	End	PSI BLAST	Verify	PMF	SeqFold	Coumpound	PDB annotation
		4	4	Score	31016	31000	2000		
1fvd	А	11	225	8.5e-72			76.29	IMMUNOGLOBULIN FAB FRAGMENT OF	
								HUMANIZED ANTIBODY 4D5, VERSION 4 1FVD 3	
1kb5	7	11	227	6.8e-72			73.99	KB5-C20 T-CELL ANTIGEN RECEPTOR; CHAIN: A, B;	COMPLEX (IMMUNOGLOBULIN/RECEPTOR) TCR
								ANTIBODY DESIRE-1; CHAIN: L, H;	VAPLHA VBETA DOMAIN; T-CELL RECEPTOR, STRAND SWITCH, FAB, ANTECT CALOCYPIC 3
									(IMMUNOGLOBULIN/RECEPTOR)
lnca		18	227	8.5e-72			73.83	HYDROLASE(O.	
								GLYCOSYL) N9 NEURAMINIDASE-NC41	
	-							(E.C.3.2.1.18) COMPLEX WITH FAB 1NCA 3	
1nfd	В	18	235	6.8e-25			77.54	NIS ALPHA-BETA T-CELL	COMPLEX
								RECEPTOR; CHAIN: A, B, C. D. H57 FAB: CHAIN: E.	(IMMUNORECEPTOR/IMMUNOGLOBU
						 -		F, G, H	(IMMUNORECEPTOR/IMMUNOGLOBU
losp	L	17	227	5.1e-67			73.22	FAB 184.1; CHAIN: L, H;	COMPLEX
								OUTER SURFACE	(IMMUNOGLOBULIN/LIPOPROTEIN)
								rkolelin A; Chalin: U;	OSFA; COMPLEA MAMINOGI OBIII INA IPOPROTEIN)
	٠								OUTER SURFACE 2 PROTEIN A
						_			COMPLEXED WITH FAB184.1,
									BORRELIA BURGDORFERI 3 STRAIN B31
ltcr	æ	18	235	5.1e-22			73.74	ALPHA, BETA T-CELL	RECEPTOR TCR, T-CELL, RECEPTOR,
		•						NECET FOR CHAIN, A, B,	SIGNAL
lvge	,,,	17	225	8.5e-72			75.01	TRI.9 FAB; CHAIN: L, H;	IMMUNOGLOBULIN TRI.9, ANTI-
									IHYROID PEROXIDASE, AUTOANTIBODY, 2

ᄺ	PDB Ch	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
	-	-			Score					IMMUNOGLOBULIN
1.≅	25c8 L		18	225	1.5e-73			74.57	IGG 5C8; CHAIN: L, H;	CATALYTIC ANTIBODY CATALYTIC ANTIBODY, FAB, RING CLOSURE REACTION
l 🛱	2mcg 1		17	226	6.8e-63			75.07	IMMUNOGLOBULIN IMMUNOGLOBULIN LAMBDA LIGHT CHAIN DIMER (MCG\$) 2MCG 3 (TRIGONAL FORM) 2MCG	
7fab	J.		81	222	1e-59			76.52	IMMUNOGLOBULIN IMMUNOGLOBULIN FAB' NEW (LAMBDA LIGHT CHAIN) 7FAB 3	
. <u>G</u>	ladq L		91	242	8.5e-57			67.94	IGG4 REA; CHAIN: A; RF- AN IGM/LAMBDA; CHAIN: H, L;	COMPLEX (IMMUNOGLOBULIN/AUTOANTIGEN) COMPLEX (IMMINOGLOBULIN/AUTOANTIGEN)
										RHEUMATOID FACTOR 2 AUTO-
laif	f		18	240	5.1e-63	0.26	0.25		ANTI-IDIOTYPIC FAB 409.5.3 (IGG2A) FAB; CHAIN: A, B, L, H	IMMUNOGLOBULIN IMMUNOGLOBULIN, C REGION, V REGION
i X	lao7 E		18	250	3.4e-27			69.03	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA-A2 HEAVY CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2
	·								RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR
rg i	1bd2 E		18	250	1.7e-38			77.88	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULN; CHAIN: B; TAX PEPTIDE;	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA A2 HEAVY CHAIN; COMPLEX (MHC/VIRAL

SEQ	PDB	Chain	Start	End	PSI	Verify	PMF	SeqFold	Coumpound	PDB annotation
NO:	a	a l	AA	AA	BLAST Score	Score	Score	Score		
			,						CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL	PEPTIDE/RECEPTOR)
						-			RECEPTOR BETA; CHAIN: E;	
11.09	1 bec		18	250	1.7e-32			76.74	14.3.D T CELL ANTIGEN RECEPTOR; 1BEC 5	RECEPTOR T CELL RECEPTOR IBEC 14
									CHAIN: NULL; 1BEC 6	
671	1bih	V	11	419	2.7e-36			90.93	HEMOLIN; CHAIN: A, B;	INSECT IMMUNITY INSECT IMMUNITY, LPS-BINDING, HOMOPHI IC ADHESION
671	19:1	1-1	18	240	6 80-63	96.0	0.13		FAR FRAGMENT: CHAIN:	COMPI EY (ANTIBODY/ANTIGEN)
	·)	2	?		3	;		L, H, J, K; VASCULAR	FAB-12; VEGF; COMPLEX
					·				ENDOTHELIAL GROWTH	(ANTIBODY/ANTIGEN), ANGIOGENIC
									FACTOR; CHAIN: V, W;	FACTOR
671	1bjm	∢	17	242	1.5e-54			67.92	LOC - LAMBDA 1 TYPE	IMMUNOGLOBULIN BENCE-JONES
									LIGHT-CHAIN DIMER;	PROTEIN; IBJM 8 BENCE JONES, ANTIBODY MILTIPLE OLIATERNARY
									7	STRUCTURES 1BJM 13
671	1bvk	4	18	119	3.4e-33	0.24	0.00		HULYSII; CHAIN: A, B, D, E; LYSOZYME; CHAIN: C,	COMPLEX (HUMANIZED ANTIBODY/HYDROLASE)
-									T.	MURAMIDASE; HUMANIŹED
										ANTIBODY, ANTIBODY COMPLEX,
										FV, ANTI-LYSOZYME, 2 COMPLEX
										(HUMANIZED ANTIBODY/HYDROLASE)
671	Ibww	A	28	118	1.7e-32	0.81	-0.05		IG KAPPA CHAIN V-I	IMMUNE SYSTEM REIV, STABILIZED
					•	,			REGION REI; CHAIN: A, B;	IMMUNOGLOBULIN FRAGMENT,
					,					BENCE-JONES 2 PROTEIN, IMMUNE SYSTEM
1/9	lcdy		254	331	5.4e-14	0.20	0.35		T-CELL SURFACE	T-CELL SURFACE GLYCOPROTEIN
					-				CHAIN: NULL;	TRANSMEMBRANE, GLYCOPROTEIN,
										T-CELL, 2 MHC, LIPOPROTEIN, T-

PDB annotation	CELL SURFACE GLYCOPROTEIN	CATALYTIC ANTIBODY CATALYTIC ANTIBODY, TERPENOID SYNTHASE, CARBOCATION, 2 CYCLIZATION Y CASCADE	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF, FGFR, IMMUNOGLOBULIN-LIKE, SIGNAL TRANSDUCTION, 2 DIMERIZATION, GROWTH FACTOR/GROWTH FACTOR RECEPTOR	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF, FGFR, IMMUNOGLOBULIN-LIKE, SIGNAL TRANSDUCTION, 2 DIMERIZATION, GROWTH FACTOR/GROWTH FACTOR	COMPLEX (ANTIBODY ANTIGEN) 1,4-BETA-N-ACETYLMURAMIDASE C; SINGLE-DOMAIN ANTIBODY, TURKEY EGG-WHITE LYSOZYME, 2 ANTIBODY-PROTEIN COMPLEX, SINGLE-CHAIN FV FRAGMENT	OELL ADHESION NCAM; NCAM, IMMUNOGLOBULIN FOLD, GLYCOPROTEIN	GROWTH FACTOR/GROWTH FACTOR C, RECEPTOR FGF2; FGFR2; H IMMUNOGLOBULIN (IG)LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS. B-TREFOIL FOLD	GROWTH FACTOR/GROWTH FACTOR
Coumpound		CATALYTIC ANTIBODY 19A4 (LIGHT CHAIN); CHAIN: L; CATALYTIC ANTIBODY 19A4 (HEAVY CHAIN); CHAIN: H;	FIBROBLAST GROWTH FACTOR 2, CHAIN: A, B; FIBROBLAST GROWTH FACTOR RECEPTOR 1; CHAIN: C, D;	FIBROBLAST GROWTH FACTOR 2; CHAIN: A, B; FIBROBLAST GROWTH FACTOR RECEPTOR 1; CHAIN: C, D;	SCFV FRAGMENT 1F9; CHAIN: A, B; TURKEY EGG-WHITE LYSOZYME C; CHAIN: X, Y;	NEURAL CELL ADHESION MOLECULE; CHAIN: A, B, C, D;	FIBROBLAST GROWTH FACTOR 2; CHAIN: A, B, C, D; FIBROBLAST GROWTH FACTOR RECEPTOR 2; CHAIN: E, F, G, H;	FIREORI AST GROWTH
SeqFold Score								
PMF Score		-0.03	-0.01	-0.03	-0.19	0.92	0.22	0.28
Verify Score		0.31	0.37	0.33	0.19	0.41	0.36	0.30
PSI BLAST Score		3.4e-63	6.8e-47	le-48	1.4e-12	5.4e-17	1.7e-45	5.1e-14
End AA		240	334	334	288	334	334	344
Start AA		20	135	135	139	159	132	265
Chain ID		,1	U	Q	∢	4	ы	ш
PDB ID		lcf8	lcvs	Icvs	1dzb	lepf	lev2	lev2
SEQ No:		129	129	671	671	671	671	671

	I-SET	ACTOR 1-SET	CCTOR I-SET	CTOR I-SET	TY I) IGE- IE- SODY,	
tation	R2; IG)LIKE G TO THE IG-LIKE	R2; IG)LIKE G TO THE IG-LIKE	RI; IG) LIKE G TO THE IG-LIKE	RI; IG) LIKE G TO THE IG-LIKE	SH AFFINI CEPSILON IN FOLD, EPTOR, IC	
PDB annotation	FGF2; FGF OBULIN (ELONGIN P WITHIN	ACTOR/GF FGF2; FGF OBULIN (ELONGIN P WITHIN	ACTOR/GR GGF1; FGF OBULIN (ELONGIN P WITHIN	ACTOR/GR GGF1; FGF OBULIN (ELONGIN P WITHIN	STEM HIC EPTOR, FC OGLOBUL IEIN, REC ROTEIN,	
	RECEPTOR FGF2; FGFR2; IMMUNOGLOBULIN (IG)LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS, B-TREFOIL FOLD	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF2; FGFR2; IMMUNOGLOBULIN (IG)LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS. B-TREFOIL FOLD	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF1; FGFR1; IMMUNOGLOBULIN (IG) LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS, B-TREFOIL FOLD	GROWTH FACTOR/GROWTH FACTOR RECEPTOR FGF1; FGFR1; IMMUNOGLOBULIN (IG) LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS, B-TREFOIL FOLD	IMMUNE SYSTEM HIGH AFFINITY IGE-FC RECEPTOR, FC(EPSILON) IGE- FC; IMMUNOGLOBULIN FOLD, GLYCOPROTEIN, RECEPTOR, IGE- BINDING 2 PROTEIN, IGE ANTIBODY, IGE-FC	
	A, B, C, OWTH ?2;	/ТН А, В, С, ОWТН К.2;	/TH A, B; /TH t I;	/TH A, B; /TH (1);	, c S S SHAIN:	ON OF IV 3
Coumpound	CHAIN: ILAST GR ECEPTOR F, G, H;	ST GROW; CHAIN: LAST GREEPTOR	ST GROW CHAIN: AST GROW ECEPTOR D;	ST GROW CHAIN: AST GROW ECEPTOR D;	NITY LOBULIN ECEPTOR IG EPSIL(EGION; C	ILOBULIN TOFA ED VERSI CD18 1FG
ပိ	FACTOR 2; CHAIN: A, B, C, D; FIBROBLAST GROWTH FACTOR RECEPTOR 2; CHAIN: E, F, G, H;	FIBROBLAST GROWTH FACTOR 2; CHAIN: A, B, C, D; FIBROBLAST GROWTH FACTOR RECEPTOR 2; CHAIN: E, F, G, H;	FIBROBLAST GROWTH FACTOR I; CHAIN: A, B; FIBROBLAST GROWTH FACTOR RECEPTOR I; CHAIN: C, D;	FIBROBLAST GROWTH FACTOR 1; CHAIN: A, B; FIBROBLAST GROWTH FACTOR RECEPTOR 1; CHAIN: C, D;	HIGH AFFINITY IMMUNOGLOBULIN EPSILON RECEPTOR CHAIN: A; IG EPSILON CHAIN C REGION; CHAIN: B, D:	IMMUNOGLOBULIN FV FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 IFGV 3 ANTIBODY H52' (HUH52-
SeqFold Score						
PMF Score		90.0	-0.08	0.16	-0.01	-0.05
Verify Score		0.40	0.40	0.62	0.35	0.29
PSI BLAST		1.4e-48	5.1e-49	5.4e-14	2.4e-14	1.4e-34
End		338	334	339	339	118
Start AA		134	135	247	159	18
Chain ID		O	U	U	A	J
PDB ID		lev2	levt	levt	1f6a	lfgv
SEQ TD		671	1/9	671	671	671

A	Chain	Start	End AA	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
\top				Score					
16.0		2	240	5 10-63	0.17	0.27		AA FV) IFGV 4	
n	1	3	2	3		1		IMMUNOGLOBULIN G1	
								(KAPPA LIGHT CHAIN) FAB' FRAGMENT 1FIG 3	
1fns	I	81	240	3.4e-63	0.36	0.12		IMMUNOGLOBULIN NMC-	IMMUNE SYSTEM VON WILLEBRAND
								4 IGG1; CHAIN: L;	FACTOR, GLYCOPROTEIN IBA
		•						IMMUNOGLOBULIN NMC-	(A:ALPHA) BINDING, 2 COMPLEX
					.,			4 IGG1; CHAIN: H; VON	(WILLEBRAND/IMMUNOGLOBULIN),
								WILLEBRAND FACTOR;	BLOOD COAGULATION TYPE 3 2B
								CHAIN: A;	VON WILLEBRAND DISEASE
1fvc	A	81	120	1.2e-32	0.65	-0.14		IMMUNOGLOBULIN FV	
								FRAGMENT OF	
								HUMANIZED ANTIBODY	
			·					4D5, VERSION 8 1FVC 3	
1fvd	¥	17	241	6.8e-61			66.39	IMMUNOGLOBULIN FAB	
								FRAGMENT OF	
								HUMANIZED ANTIBODY	
7								4D5, VERSION 4 1FVD 3	
lhng	A	159	338	1.9e-18	0.08	-0.14		TLYMPHOCYTE	
_								AUTESION	
-								GLYCOPROTEIN CD2	
lion.		<u>~</u>	126	8 50-34	0.46	100		PANTINOGI OBLILIN	
_	1	2	2		?	5		IMMUNOGLOBULIN M	
								(IG-M) FV FRAGMENT	
								IIGM 3	
litb	В	139	420	1.9e-14			69.14	INTERLEUKIN-1 BETA;	COMPLEX
								CHAIN: A; TYPE 1	(IMMUNOGLOBULIN/RECEPTOR)
								INTERLEUKIN-1	IMMUNOGLOBULIN FOLD,
								RECEPTOR; CHAIN: B;	TRANSMEMBRANE, GLYCOPROTEIN,
									RECEPTOR, 2 SIGNAL, COMPLEX (IMMUNOGLOBULIN/RECEPTOR)
£1	В	154	362	1.9e-14	0.34	0.09		INTERLEUKIN-1 BETA;	COMPLEX

PDB Ch	Chain ID	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
								CHAIN: A; TYPE 1 INTERLEUKIN-1 RECEPTOR; CHAIN: B;	(IMMUNOGLOBULINRECEPTOR) IMMUNOGLOBULIN FOLD, TRANSMEMBRANE, GLYCOPROTEIN, RECEPTOR, 2 SIGNAL, COMPLEX (IMMUNOGLOBULINRECEPTOR)
Imcp L		19	240	8.5e-63	0.44	90:0		IMMUNOGLOBULIN IMMUNOGLOBULIN FAB FRAGMENT (MCPC\$603) IMCP 4	
		255	331	8.je-15	0.44	0.07		TITIN; CHAIN: NULL;	MUSCLE PROTEIN CONNECTIN, NEXTM5; CELL ADHESION, GLYCOPROTEIN, TRANSMEMBRANE, REPEAT, BRAIN, 2 IMMUNOGLOBULIN FOLD, ALTERNATIVE SPLICING, SIGNAL, 3 MUSCLE PROTEIN
φ	j	18	251	5.1e-35			76.63	NIS ALPHA-BETA T-CELL RECEPTOR; CHAIN: A, B, C, D; HS7 FAB; CHAIN: E, F, G, H	COMPLEX (IMMUNORECEPTOR/IMMUNOGLOBU LIN) COMPLEX (IMMUNORECEPTOR/IMMUNOGLOBU LIN)
lqac A		19	611	1.7e-32	0.55	-0.05		IMMUNOGLOBULIN LIGHT CHAIN VARIABLE DOMAIN; CHAIN: A, B;	IMMUNE SYSTEM BETA BARREL IMMUNOGLOBULIN VL DOMAIN DIMER, FLIPPED DOMAIN 2 DIMER
lqnz L		20	119	1.7e-31	0.37	9.06		0.5B ANTIBODY (LIGHT CHAIN); CHAIN: L; 0.5B ANTIBODY (HEAVY CHAIN); CHAIN: H; GP120; CHAIN: P;	ANTIBODY ANTIBODY, V3 PEPTIDE, BINDING SITE
ח		61	240	3.4e-63	0.42	0.03		MONOCLONAL ANTIBODY 3A2; CHAIN: H, L;	MONOCLONAL ANTIBODY MONOCLONAL ANTIBODY, FAB- FRAGMENT, REPRODUCTION
В		18	251	1.7e-33			70.09	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN,

SEQ ID NO:	PDB ID	Chain ID	Start	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
										SIGNAL
1.69	Itnm		255	331	2.7e-15	0.55	90.0		MUSCLE PROTEIN TITIN MODULE M5 (CONNECTIN) 1TNM 3 (NMR, MINIMIZED AVERAGE STRUCTURE) 1TNM 4 1TNM 58	
129	lwej	П	18	240	8.5e-63	0.34	0.05		E8 ANTIBODY; CHAIN: L, H; CYTOCHROME C; CHAIN: F;	COMPLEX (ANTIBODY/ELECTRON TRANSPORT) FAB E8; CYT C, ANTIGEN; IMMUNOGLOBULIN, IGGI KAPPA, FAB FRAGMENT, HORSE 2 CYTOCHROME C, COMPLEX (ANTIBODY/ELECTRON TRANSPORT)
129	1wio	Ą	23	365	1.4e-25			78.38	T-CELL SURFACE GLYCOPROTEIN CD4; CHAIN: A, B;	GLYCOPROTEIN CD4; IMMUNOGLOBULIN FOLD, TRANSMEMBRANE, GLYCOPROTEIN, T-CELL, 2 MHC LIPOPROTEIN, POLYMORPHISM
671	lwit		257	331	1.1e-14	0.51	-0.15		TWITCHIN 18TH IGSF MODULE; CHAIN: NULL;	MUSCLE PROTEIN IMMUNOGLOBULIN SUPERFAMILY, I SET, MUSCLE PROTEIN
671	Iwtl	A	18	119	6.8e-32	0.51	-0.11		IMMUNOGLOBULIN WAT, A VARIABLE DOMAIN FROM IMMUNOGLOBULIN LIGHT-CHAIN IWTL 3 (BENCE-JONES PROTEIN)	
119	25c8	1	20	240	1.7e-63	0:39	0.04		IGG 5C8; CHAIN: L, H;	CATALYTIC ANTIBODY CATALYTIC ANTIBODY, FAB, RING CLOSURE REACTION
671	2dli	А	159	333	1.1e-14	0.05	-0.06		MHC CLASS I NK CELL RECEPTOR PRECURSOR; CHAIN: A;	IMMUNE SYSTEM PS8 NATURAL KILLER CELL RECEPTOR; KIR, NATURAL KILLER RECEPTOR,

PDB annotation	INHIBITORY RECEPTOR, 2 IMMUNOGLOBULIN				CELL ADHESION NCAM DOMAIN 1; CELL ADHESION, GLYCOPROTEIN, HEPARIN-BINDING, GPI-ANCHOR, 2 NEURAL ADHESION MOLECULE, IMMUNOGLOBULIN FOLD, SIGNAL	CELL ADHESION PROTEIN NCAM MODULE 2; CELL ADHESION, GLYCOPROTEIN, HEPARIN-BINDING,
Coumpound		IMMUNOGLOBULIN FAB FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4	IMUNOGLOBULIN IMMUNOGLOBULIN VL DOMAIN (VARIABLE DOMAIN OF KAPPA ZIMN 3 LIGHT CHAIN) OF MCPC603 MUTANT IN WHICH ZIMN 4 COMPLEMENTARITY- DETERMINING REGION I HAS BEEN REPLACED BY ZIMN 5 THAT FROM MOPC167 ZIMN 6	IMMUNOGLOBULIN IMMUNOGLOBULIN LAMBDA LIGHT CHAIN DIMER (MCG\$) 2MCG 3 (TRIGONAL FORM) 2MCG	NEURAL CELL ADHESION MOLECULE; CHAIN: NULL;	NEURAL CELL ADHESION MOLECULE, LARGE ISOFORM; CHAIN: A;
SeqFold Score				68.68		
PMF Score		0.42	0.28		0.18	0.34
Verify Score		0.42	69.0		1.06	0.51
PSI BLAST Score		3.4e-63	16-31	5.1e-56	5.4e-15	5.4e-15
End AA		240	119	242	338	331
Start AA		81	19	17	255	255
Chain ID		ı		-		А
PDB ID		2fgw	2imn	2тсд	2ncm	Зпст
SEQ ID NO:		671	671	129	671	671

PDB annotation	GPI-ANCHOR, 2 NEURAL ADHESION MOLECULE, IMMUNOGLOBULIN FOLD, HOMOPHILIC 3 BINDING, CELL ADHESION PROTEIN		IMMUNOGLOBULIN IMMUNOGLOBULIN	IMMUNOGLOBULIN IMMUNOGLOBULIN, C'REGION, V REGION		COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA A2 HEAVY CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR)	RECEPTOR T CELL RECEPTOR IBEC 14	INSECT IMMUNITY INSECT IMMUNITY, LPS-BINDING,
Coumpound		IMMUNOGLOBULIN IMMUNOGLOBULIN FAB' NEW (LAMBDA LIGHT CHAIN) 7FAB 3	2E8 (IGGI=KAPPA=) ANTIBODY; CHAIN: L, H, M, P;	ANTI-IDIOTYPIC FAB 409.5.3 (IGG2A) FAB; CHAIN: A, B, L, H	IMMUNOGLOBULIN FAB' FRAGMENT OF MONOCLONAL ANTIBODY B72.3 1BBJ 3 (MURINE/HUMAN CHIMERA) 1BBJ 4	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	14.3.D T CELL ANTIGEN RECEPTOR; IBEC 5 CHAIN: NULL; IBEC 6	HEMOLIN; CHAIN: A, B;
SeqFold Score		68.11	73.68	73.37	72.88	72.64	75.74	102.10
PMF Score				i				
Verify Score								
PSI BLAST Score		1.7e-53	1.7e-68	5.1e-72	5.1e-69	1.2e-22	1.7e-21	8.5e-16
End		238	722	227	222	234	234	403
Start		18	18	17	17	8	18	17
Chain ID		J	T	7	J	m		A
PDB ID		7fab	12e8	laif	1 b bj	1bd2	1pec	1bih
SEQ ID NO:		671	672	672	672	672	672	672

PDB annotation	HOMOPHILIC ADHESION	ANTIBODY, CD\$2								(IMMUNOGLOBULIN/RECEPTOR) TCR	RECEPTOR, STRAND SWITCH, FAB,	ANTICLONOTYPIC, 2 (IMMUNOGLOBULIN/RECEPTOR)					COMPLEX	(IMMUNORECEPTOR/IMMUNOGLOBU	LIN) COMPLEX	(IMMUNORECEPTOR/IMMUNOGLOBU	COMPLEX	(IMMUNOGLOBULIN/LIPOPROTEIN)	OSPA; COMPLEX
Coumpound		CAMPATH-1H:LIGHT CHAIN; CHAIN: L; CAMPATH-1H:HEAVY CHAIN; CHAIN: H; PEPTIDE ANTIGEN;	CHAIN: P;	IMMUNOGLOBULIN	(KAPPA LIGHT CHAIN)	FAB FRAUMENT IFIG 3	IMMUNOGLOBULIN FAB FRAGMENT OF	HUMANIZED ANTIBODY	4D5, VERSION 4 1FVD 3	KB5-C20 T-CELL ANTIGEN RECEPTOR; CHAIN: A, B; ANTIRODY DESIRE-1:	CHAIN: L, H;		HYDROLASE(O-	GLYCOSYL) N9	(E.C.3.2.1.18) COMPLEX	WITH FAB INCA 3	NIS ALPHA-BETA T-CELL	RECEPTOR; CHAIN: A, B,	C, D; H57 FAB; CHAIN: E,	F, G, H	FAB 184.1; CHAIN: L. H.	OUTER SURFACE	PROTEIN A; CHAIN: O;
SeqFold Score		72.55		73.31			76.29			73.99			73.83				77.54				73.22		
PMF Score																							
Verify Score																							
PSI BLAST Score		5.1e-72		3.4e-72			8.5e-72			6.8e-72			8.5e-72				6.8e-25				5.1e-67		
End AA		222		227			225			227			227				235				227		
Start AA		17	4	18			17			17			18				18				17		
Chain ID		∟		7			∢			.i			7				В				T.		
PDB ID		1ce1		lfig			1 fvd			1kb5			lnca				lnfd				losp		
SEQ ID NO:		672		672			672			672			672				672				672		

PDB annotation	(IMMUNOGLOBULINILIPOPROTEIN), OUTER SURFACE 2 PROTEIN A COMPLEXED WITH FAB184.1, BORRELIA BURGDORFERI 3 STRAIN B31	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL	IMMUNOGLOBULIN TRI.9, ANTI- THYROID PEROXIDASE, AUTOANTIBODY, 2 IMMUNOGLOBULIN	CATALYTIC ANTIBODY CATALYTIC ANTIBODY, FAB, RING CLOSURE REACTION			COMPLEX (IMMUNOGLOBULIN/AUTOANTIGEN) COMPLEX (IMMUNOGLOBULIN/AUTOANTIGEN), RHEUMATOID FACTOR 2 AUTO- ANTIBODY COMPLEX	IMMUNOGLOBULIN IMMUNOGLOBULIN, C REGION, V REGION
Coumpound		ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	TR1.9 FAB; CHAIN: L, H;	IGG 5C8; CHAIN: L, H;	IMMUNOGLOBULIN IMMUNOGLOBULIN LAMBDA LIGHT CHAIN DIMER (/MCG\$) 2MCG 3 (TRIGONAL FORM) 2MCG	IMMUNOGLOBULIN IMMUNOGLOBULIN FAB' NEW (LAMBDA LIGHT CHAIN) 7FAB 3	IGGA REA; CHAIN: A; RF- AN IGMLAMBDA; CHAIN: H, L;	ANTI-IDIOTYPIC FAB 409.5.3 (IGG2A) FAB; CHAIN: A, B, L, H
SeqFold Score		73.74	75.01	74.57	75.07	76.52	67.94	
PMF Score								0.25
Verify Score								0.26
PSI BLAST Score		5.1e-22	8.5e-72	1.5e-73	6.8e-63	1e-59	8.5e-57	5.1e-63
End AA		235	225	225	226	222	242	240
Start AA		18	17	18	17	18	61	18
Chain ID		В	7	1		L	7	T
PDB ID		ltcr	lvge	25c8	2тсд	7fab	ladq	laif
SEQ ID NO:		672	672	672	672	672	672	672

PDB annotation	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA-A2 HEAVY CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA A2 HEAVY CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR)	RECEPTOR T CELL RECEPTOR IBEC 14	INSECT IMMUNITY INSECT IMMUNITY, LPS-BINDING, HOMOPHILIC ADHESION	COMPLEX (ANTIBODY/ANTIGEN) FAB-12; VEGF; COMPLEX (ANTIBODY/ANTIGEN), ANGIOGENIC FACTOR	IMMUNOGLOBULIN BENCE-JONES PROTEIN; 1BJM 8 BENCE JONES, ANTIBODY, MULTIPLE QUATERNARY STRUCTURES 1BJM 13	COMPLEX (HUMANIZED ANTIBODY/HYDROLASE) MURAMIDASE; HUMANIZED ANTIBODY, ANTIBODY COMPLEX,
	COMPLE PEPTIDE CHAIN; C RECEPTC COMPLE PEPTIDE	COMPLE PEPTIDE CHAIN; C PEPTIDE	RECEPTO 14	INSECT I IMMUNIT HOMOPH	COMPLE FAB-12; V (ANTIBO FACTOR	IMMUNO PROTEIN ANTIBOI STRUCTI	COMPLE ANTIBOL MURAMI ANTIBOL
Coumpound	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	14.3.D T CELL ANTIGEN RECEPTOR; 1BEC 5 CHAIN: NULL; 1BEC 6	HEMOLIN; CHAIN: A, B;	FAB FRAGMENT; CHAIN: I, H, J, K; VASCULAR ENDOTHELIAL GROWTH FACTOR; CHAIN: V, W;	LOC - LAMBDA I TYPE LIGHT-CHAIN DIMER; 1BJM 6 CHAIN: A, B; 1BJM 7	HULYSI I; CHAIN: A, B, D, E; LYSOZYME; CHAIN: C, F;
SeqFold Score	69.03	77.88	76.74	90.93		67.92	
PMF Score					0.13		0.00
Verify Score					0.26		0.24
PSI BLAST Score	3.4e-27	1.7e-38	1.7e-32	2.7e-36	6.8e-63	1.5e-54	3.4e-33
End	250	250	250	419	240	242	119
Start	80	81	8I	17	18	17	18
Chain	п	ш		A	7	A	A
PDB CI	1ao7	1bd2	1bec	1bih	16j1	1bjm	1bvk
SEQ ID NO:	672	672	672	672	672	672	672

Coumpound PDB annotation	FV, ANTI-LYSOZYME, 2 COMPLEX (HUMANIZED ANTIBODY/HYDROLASE)	IG KAPPA CHAIN V-I IMMUNE SYSTEM REIV, STABILIZED REGION REI; CHAIN: A, B; BENCE-JONES 2 PROTEIN, IMMUNE SYSTEM	T-CELL SURFACE T-CELL SURFACE GLYCOPROTEIN GLYCOPROTEIN CD4; IMMUNOGLOBULIN FOLD, TRANSMEMBRANE, GLYCOPROTEIN, T-CELL, 2 MHC, LIPOPROTEIN, T-CELL, 2 WHC, LIPOPROTEIN, T-CELL SURFACE GLYCOPROTEIN	CATALYTIC ANTIBODY CATALYTIC ANTIBODY CATALYTIC 19A4 (LIGHT CHAIN); ANTIBODY, TERPENOID SYNTHASE, CARBOCATION, 2 CYCLIZATION ANTIBODY 19A4 (HEAVY CASCADE CHAIN); CHAIN: H;	FIBROBLAST GROWTH FACTOR 2; CHAIN: A, B; FECEPTOR FGF, FGFR, FIBROBLAST GROWTH FACTOR RECEPTOR 1; FACTOR RECEPTOR 1; FACTOR RECEPTOR 1; GROWTH FACTOR/GROWTH FACTOR RECEPTOR	FIBROBLAST GROWTH GROWTH FACTOR/GROWTH FACTOR FACTOR 2; CHAIN: A, B; RECEPTOR FGF, FGFR, IMMUNOGLOBULIN-LIKE, SIGNAL IMMUNOGLOBULIN-LIKE, SIGNAL TACTOR RECEPTOR 1; GROWTH FACTOR/GROWTH FACTOR RECEPTOR	SCEV FRAGMENT 1F9; COMPLEX (ANTIBODY ANTIGEN) 1,4- CHAIN: A, B; TURKEY BETA-N-ACETYLMURAMIDASE C; EGG-WHITE LYSOZYME SINGLE-DOMAIN ANTIBODY, EGG-WHITE LYSOZYME BETA-N-ACETYLMURAMIDASE C; EGG-WHITE LYSOZYME BETA-N-ACETYLMURAMIDASE C; EGG-WHITE LYSOZYME BETA-N-ACETYLMURAMIN ANTIBODY, EGG-WHITE LYSOZYME BETA-N-ACETYLMURAMIN ANTIBOTY ANT
		IG KAPPA REGION R	T-CELL SURFACE GLYCOPROTEIN (CHAIN: NULL;	CATALYT 19A4 (LIG CHAIN: L; ANTIBOD CHAIN); C	FIBROBLAST FACTOR 2; C FIBROBLAST FACTOR REC CHAIN: C, D;	FIBROBLAST FACTOR 2; C FIBROBLAST FACTOR REC CHAIN: C, D;	CHAIN: A, B; TC EGG-WHITE LY
SeqFold Score							
PMF Score		-0.05	0.35	-0.03	-0.01	-0.03	-0.19
Verify Score		0.81	0.20	0.31	0.37	0.33	0.19
PSI BLAST Score		1.7e-32	5.4e-14	3.4e-63	6.8e-47	1e-48	1.4e-12
End		118	331	240	334	334	288
Start		81	254	20	135	135	139
Chain ID		4		1	ပ	Q	¥
PDB ED		1bww	1cdy	10f8	lcvs	Icvs	1dzb
SEQ ID NO:		672	672	672	672	672	672

Coumpound PDB annotation	SINGLE-CHAIN FV FRAGMENT	NEURAL CELL ADHESION CELL ADHESION NCAM; NCAM, MOLECULE; CHAIN: A, B, IMMUNOGLOBULIN FOLD, C, D:	OBLAST GROWTH TOR 2; CHAIN: A, B, C, BROBLAST GROWTH OR RECEPTOR 2; N: E, F, G, H;	FIBROBLAST GROWTH GROWTH FACTOR/GROWTH FACTOR FACTOR 2, CHAIN: A, B, C, D; FIBROBLAST GROWTH IMMUNOGLOBULIN (IG)LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS. B-TREFOLL FOLD	FIBROBLAST GROWTH FACTOR/GROWTH FACTOR FACTOR 2; CHAIN: A, B, C, RECEPTOR FGF2; FGFR2; D; FIBROBLAST GROWTH MMUNOGLOBULIN (IG)LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS. B-TREFOIL FOLD	FIBROBLAST GROWTH FACTOR/GROWTH FACTOR FACTOR 1; CHAIN: A, B; FIBROBLAST GROWTH FACTOR FGF1; FGFR1; FIBROBLAST GROWTH MMUNOGLOBULIN (IG) LIKE DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS, B-TREFOIL FOLD	FIBROBLAST GROWTH GROWTH FACTOR/GROWTH FACTOR FACTOR 1; CHAIN: A, B; RECEPTOR FGF1; FGFR1; FGFR1; FIBROBLAST GROWTH FACTOR FACTOR RECEPTOR 1; DOMAINS BELONGING TO THE I-SET 2 SUBGROUP WITHIN IG-LIKE DOMAINS. B-TREFOIL FOLD
P .		NEU MOL	FIBE FAC D; F. CHA	FIBH FAC D; FI FAC CHA	FIBE FAC D; FI FAC CHA	FIBR FAC FIBR FAC CHA	FIBR FAC FIBR CHA
SeqFold Score							
PMF		0.92	0.22	0.28	0.06	-0.08	0.16
Verify Score		0.41	0.36	0.30	0.40	0.40	0.62
PSI BLAST Score		5.4e-17	1.7e-45	5.1e-14	1.4e-48	5.1e-49	5.4e-14
End		334	334	344	338	334	339
Start AA		159	132	265	134	135	247
Chain ID		A	ш	ы	Ö	ပ	O
PDB ID		lepf	lev2	lev2	lev2	levt	levt
SEQ ID NO:		229	672	672	672	672	672

Coumpound PDB annotation	HIGH AFFINITY IMMUNOGLOBULIN EPSILON RECEPTOR FC; IMMUNOGLOBULIN FOLD, FC; IMMUNOGLOBULIN FOLD, CHAIN: A; IG EPSILON GLYCOPROTEIN, RECEPTOR, IGE- CHAIN C REGION; CHAIN: BINDING 2 PROTEIN, IGE ANTIBODY, IGE-FC	[IL	BULIN BULIN GI IT CHAIN)	IMMUNOGLOBULIN NMC- FACTOR, GLYCOPROTEIN IBA IMMUNOGLOBULIN NMC- (A:ALPHA) BINDING, 2 COMPLEX 4 IGG1; CHAIN: H; VON (WILLEBRAND/IMMUNOGLOBULIN), BLOOD COAGULATION TYPE 3 2B VON WILLEBRAND DISEASE	 	BULIN FAB DF ANTIBODY 14 IFVD 3	TE
	HIGH AFFINITY IMMUNOGLOBULIN EPSILON RECEPTOR CHAIN: A; IG EPSILON CHAIN C REGION; CHA	IMMUNOGLOBULIN FV FRAGMENT OF A HUMANIZED VERSION O THE ANTI-CD18 1FGV 3 ANTIBODY 'H52' (HUH52- AA FV) 1FGV 4	IMMUNOGLOBULIN IMMUNOGLOBULIN GI (KAPPA LIGHT CHAIN) FAB' FRAGMENT 1FIG 3	IMMUNOGLOBULIN NM 4 IGG1; CHAIN: L; IMMUNOGLOBULIN NM 4 IGG1; CHAIN: H; VON WILLEBRAND FACTOR; CHAIN: A:	IMMUNOGLOBULIN FV FRAGMENT OF HUMANIZED ANTIBODY 4D5, VERSION 8 1FVC 3	IMMUNOGLOBULIN FAB FRAGMENT OF HUMANIZED ANTIBODY 4D5: VERSION 4 1FVD 3	TLYMPHOCYTE ADHESION GI VCORPOTEIN CD2
SeqFold Score						66.39	
PMF	-0.01	-0.05	0.27	0.12	-0.14		-0.14
Verify Score	0.35	0.29	0.17	0.36	0.65		0.08
PSI BLAST Score	2,4e-14	1.4e-34	5.1e-63	3.4e-63	1.2e-32	6.8e-61	1.9e-18
End	339	118	240	240	120	241	338
Start	159	18	20	18	18	17	159
Chain	4	1	7	7	V	V	Ą
PDB ID	1f6a	Ifgv	1fig	Ifns	lfvc	1fvd	Ihng
SEQ ID NO:	672	672	672	672	672	672	672

SEQ	PDB et	Chain ID	Start AA	End	PSI BLAST	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
672 672	ligm 1	ļ.	18	126	8.5e-34	0.46	-0.01		IMMUNOGLOBULIN IMMUNOGLOBULIN M (IG-M) FV FRAGMENT	
672	litb	Ф	139	420	1.9e-14			69.14	NTERLEUKIN-1 BETA; CHAIN: A; TYPE 1 INTERLEUKIN-1 RECEPTOR; CHAIN: B;	COMPLEX (IMMUNOGLOBULIN/RECEPTOR) IMMUNOGLOBULIN FOLD, TRANSMEMBRANE, GLYCOPROTEIN, RECEPTOR, 2 SIGNAL, COMPLEX (IMMINOGLOBILI IN/RECEPTOR)
672	Titb	В	154	362	1.9e-14	0.34	0.09		INTERLEUKIN-1 BETA; CHAIN: A; TYPE 1 INTERLEUKIN-1 RECEPTOR; CHAIN: B;	COMPLEX (IMMUNOGLOBULINRECEPTOR) IMMUNOGLOBULIN FOLD, TRANSMEMBRANE, GLYCOPROTEIN, RECEPTOR, 2 SIGNAL, COMPLEX (IMMUNOGLOBULINRECEPTOR)
672	Ітср	7	19	240	8.5e-63	0.44	0.06		IMMUNOGLOBULIN IMMUNOGLOBULIN FAB FRAGMENT (MC/PC\$603) IMCP 4	
672	Inct		255	331	8.16-15	0.44	0.07		TITIN; CHAIN: NULL;	MUSCLE PROTEIN CONNECTIN, NEXTMS; CELL ADHESION, GLYCOPROTEIN, TRANSMEMBRANE, REPEAT, BRAIN, 2 IMMUNOGLOBULIN FOLD, ALTERNATIVE SPLICING, SIGNAL, 3 MUSCLE PROTEIN
672	Infd	В	18	251	5.1e-35			76.63	NIS ALPHA-BETA T-CELL RECEPTOR; CHAIN: A, B, C, D; H57 FAB; CHAIN: E, F, G, H	COMPLEX (IMMUNORECEPTOR/IMMUNOGLOBU LIN) COMPLEX (IMMUNORECEPTOR/IMMUNOGLOBU LIN)
672	lqac	V	61	611	1.7e-32	0.55	-0.05		IMMUNOGLOBULIN LIGHT CHAIN VARIABLE	IMMUNE SYSTEM BETA BARREL IMMUNOGLOBULIN VL DOMAIN

PDB annotation	DIMER, FLIPPED DOMAIN 2 DIMER		MONOCLONAL ANTIBODY MONOCLONAL ANTIBODY, FAB- FRAGMENT, REPRODUCTION	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL		COMPLEX (ANTIBODY/ELECTRON TRANSPORT) FAB E8; CYT C, ANTIGEN, IMMUNOGLOBULIN, IGG1 KAPPA, FAB FRAGMENT, HORSE 2 CYTOCHROME C, COMPLEX (ANTIBODY/ELECTRON TRANSPORT)	GLYCOPROTEIN CD4; IMMUNOGLOBULIN FOLD, TRANSMEMBRANE, GLYCOPROTEIN, T-CELL, 2 MHC LIPOPROTEIN, POLYMORPHISM	MUSCLE PROTEIN IMMUNOGLOBULIN SUPERFAMILY, I SET, MUSCLE PROTEIN	-
Coumpound	DOMAIN; CHAIN: A, B;	0.5B ANTIBODY (LIGHT CHAIN); CHAIN: L; 0.5B ANTIBODY (HEAVY CHAIN); CHAIN: H; GP120; CHAIN: P.	MONOCLONAL ANTIBODY 3A2; CHAIN: H, L:	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	MUSCLE PROTEIN TITIN MODULE MS (CONNECTIN) 1TNM 3 (NMR, MINIMIZED AVERAGE STRUCTURE)	ES ANTIBODY; CHAIN: L, H; CYTOCHROME C; CHAIN: F;	T-CELL SURFACE GLYCOPROTEIN CD4; CHAIN: A, B;	TWITCHIN 18TH IGSF MODULE; CHAIN: NULL;	IMMUNOGLOBULIN WAT,
SeqFold Score				70.09			78.38		
PMF Score		90.0	0.03		90.0	0.05		-0.15	-0.11
Verify Score		0.37	0.42		0.55	0.34		0.51	0.51
PSI BLAST Score		1.7e-31	3.4e-63	1.7e-33	2.7e-15	8.5e-63	1.4e-25	1.1e-14	6.8e-32
End AA		119	240	251	331	240	365	331	119
Start AA		20	19	18	255	81	23	. 257	81
Chain ID		7	H	В		ı	٧		¥
PDB ID		Iqnz	1sbs	ltcr	1tnm	lwej	lwio	Iwit	lwti
SEQ ID NO:		672	672	672	672	672	672	672	672

otation		DY CATALYTIC	8 NATURAL TOR; KIR, ECEPTOR, OR, 2			
PDB annotation		CATALYTIC ANTIBODY CATALYTIC ANTIBODY, FAB, RING CLOSURE REACTION	MMUNE SYSTEM PS8 NATURAL KILLER CELL RECEPTOR; KIR, NATURAL KILLER RECEPTOR, INHIBITORY RECEPTOR, IMMUNOGLOBULIN			
Coumpound	FROM IMMUNOGLOBULIN LIGHT-CHAIN 1WTL 3 (BENCE-JONES PROTEIN) 1WTL 4	IGG 5C8; CHAIN: L, H;	MHC CLASS I NK CELL RECEPTOR PRECURSOR; CHAIN: A;	IMMUNOGLOBULIN FAB FRAGMENT OF A HUMANIZED VERSION OF THE ANTI-CD18 2FGW 3 ANTIBODY 'H52' (HUH52- OZ FAB) 2FGW 4	IMUNOGLOBULIN IMMUNOGLOBULIN VL DOMAIN (VARIABLE DOMAIN OF KAPPA 2IMN 3 LIGHT CHAIN) OF MCPC603 MUTANT IN WHICH 2IMN 4 COMPLEMENTARITY- DETERMINING REGION I HAS BEEN REPLACED BY 2IMN 5 THAT FROM MOPC167 2IMN 6	IMMUNOGLOBULIN IMMUNOGLOBULIN I AMBDA I ICHT CHARI
SeqFold Score						89.89
PMF Score		0.04	-0.06	0.42	0.28	
Verify Score		0.39	0.05	0.42	69.0	
PSI BLAST Score		1.7e-63	1.1e-14	3.4e-63	16-31	5.1e-56
End AA		240	333	240	119	242
Start AA		20	159	18	61	17
Chain ID		Т	٧	1		1
PDB ID		25c8	2dli	2fgw	2imn	2mcg
SEQ ID NO:		229	672	672	672	219

PDB annotation		CELL ADHESION NCAM DOMAIN 1; CELL ADHESION, GLYCOPROTEIN, HEPARIN-BINDING, GPI-ANCHOR, 2 NEURAL ADHESION MOLECULE, MMUNOGLOBULIN FOLD, SIGNAL	CELL ADHESION PROTEIN NCAM MODULE 2; CELL ADHESION, GLYCOPROTEIN, HEPARIN-BINDING, GPI-ANCHOR, 2 NEURAL ADHESION MOLECULE, IMMUNOGLOBULIN FOLD, HOMOPHILIC 3 BINDING, CELL ADHESION PROTEIN		COMPLEX (GTP-BINDING/TRANSDUCER) BETA!, TRANSDUCIN BETA SUBUNIT; GAMMA!, TRANSDUCIN GAMMA SUBUNIT; COMPLEX (GTP-BINDING/TRANSDUCER), G PROTEIN, HETEROTRIMER 2 SIGNAL TRANSDUCTION	
		CELL AD CELL AD HEPARIN NEURAL IMMUNO	CELL AD MODULE GL YCOPI GPI-ANCI MOLECU FOLD, HC		COMPLEX (GTP-BINDING/TRANS TRANSDUCIN BE GAMMA1, TRANS SUBUNIT; COMPI BINDING/TRANS HETEROTRIMER TRANSDUCTION	
Coumpound	DIMER (MCGS) 2MCG 3 (TRIGONAL FORM) 2MCG 4	NEURAL CELL ADHESION MOLECULE; CHAIN: NULL;	NEURAL CELL ADHESION MOLECULE, LARGE ISOFORM; CHAIN: A;	IMMUNOGLOBULIN IMMUNOGLOBULIN FAB' NEW (LAMBDA LIGHT CHAIN) 7FAB 3	GT-ALPHA/GI-ALPHA CHIMERA; CHAIN: A; GT- BETA; CHAIN: B; GT- GAMMA; CHAIN: G;	GTP-BINDING PROTEIN TRANSDUCIN-ALPHA (GT- ALPHA-GDP-ALF, T- ALPHA-GDP-ALF) 1TAD 3 COMPLEXED WITH GDP AND ALUMINUM
SeqFold Score				68.11	262.16	271.77
PMF Score		0.18	0.34			
Verify Score		1.06	0.51			
PSI BLAST Score		5.4e-15	5.4e-15	1.7e-53	6.8e- 100	8.5e-95
End		338	331	238	264	264
Start AA		255	255	18	9	27
Chain ID			4	7	∢	4
PDB ID		2ncm	3пст	7fab	lgot	ltad
SEQ ID NO:		672	672	672	678	678

SEQ ID NO:	PDB ID	Chain ID	Start AA	End	PSI BLAST Score	Verify Score	PMF Score	SeqFold Score	Coumpound	PDB annotation
									FLUORIDE 1TAD 4	
619	lac6	4	21	123	8.5e-33			67.32	T-CELL RECEPTOR ALPHA; CHAIN: A, B;	RECEPTOR RECEPTOR, V ALPHA DOMAIN, SITE-DIRECTED MUTAGENESIS, 2 THREE- DIMENSIONAL STRUCTURE, GI VCOPROTEN SIGNAL
629	lac6	А	21	136	8.5e-33	0.45	1.00		T-CELL RECEPTOR ALPHA; CHAIN: A, B;	RECEPTOR RECEPTOR, VALPHA DOMAIN, SITE-DIRECTED MUTAGENESIS, 2 THREE- DIMENSIONAL STRUCTURE, GLYCOPROTEIN, SIGNAL
679	1ao7	Q	21	132				51.63	HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA-A2 HEAVY CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR
679	1ao7	Ω	23	136	3,4e-31	-0.07	0.89		HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA-42 HEAVY CHAIN; CLASS I MHC, T-CELL RECEPTOR, VIRAL PEPTIDE, 2 COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR
629	1588	A	21	136	1.5e-34	0.17	0.53		T CELL RECEPTOR V. ALPHA DOMAIN; CHAIN: A, B;	T CELL RECEPTOR TCR; T CELL RECEPTOR, MHC CLASS I, HUMAN IMMUNODEFICIENCY VIRUS, 2 MOLECULAR RECOGNITION
629	1bd2	О	21	136	1.7e-34	0.40	0.89		HLA-A 0201; CHAIN: A; BETA-2 MICROGLOBULIN;	COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR) HLA A2 HEAVY

PDB annotation		CHAIN; COMPLEX (MHC/VIRAL PEPTIDE/RECEPTOR)	IMMUNE SYSTEM IMMUNOGLOBULIN, IMMUNORECEPTOR, IMMUNE SYSTEM		COMPLEX
Coumpound		CHAIN: B; TAX PEPTIDE; CHAIN: C; T CELL RECEPTOR ALPHA; CHAIN: D; T CELL RECEPTOR BETA; CHAIN: E;	ALPHA-BETA T CELL RECEPTOR (TCR) (D10); CHAIN: A;	T-CELL RECEPTOR D10 (ALPHA CHAIN); CHAIN: A, E; T-CELL RECEPTOR D10 (BETA CHAIN); CHAIN: B, F; MHC I-AK A CHAIN (ALPHA CHAIN); CHAIN (ALPHA CHAIN); CHAIN: C, G; MHC I-AK B CHAIN (BETA CHAIN); CHAIN: P, Q; HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DR CHAIN: A; HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DR. CHAIN: B; HEMAGGLUTININ HAI PEPTIDE CHAIN; CHAIN: B; HEMAGGLUTININ HAI PEPTIDE CHAIN; CHAIN: C; T-CELL RECEPTOR ALPHA CHAIN; CHAIN: D; T-CELL RECEPTOR ALPHA CHAIN; CHAIN: D; T-CELL RECEPTOR CHAIN; CHAIN: B;	KB5-C20 T-CELL ANTIGEN
SeqFold	Score				
PMF	Score		0.90	1.00	0.13
Verify	Score		0.18	0.30	0.24
PSI BIACT	Score		3.4e-36		8.5e-37
End	AA A		136	136	139
Start	AA		16	22 21	21
Chain	3		4	A D	A
PDB	3		1bwm	149k	1kb5
SEQ	βŞ		619	679	679

PDB annotation	(IMMUNOGLOBULIN/RECEPTOR) TCR VAPLHA VBETA DOMAIN; T-CELL RECEPTOR, STRAND SWITCH, FAB, ANTICLONOTYPIC, 2 (IMMUNOGLOBULIN/RECEPTOR)	IMMUNE SYSTEM HUMAN TCR/PEPTIDE/MHC COMPLEX, HLA- A2, HTLV-1, TAX, TCR, T2 CELL RECEPTOR, IMMUNE SYSTEM	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL	RECEPTOR TCR; T-CELL, RECEPTOR, TRANSMEMBRANE, GLYCOPROTEIN, SIGNAL		DESIGNED HELICAL BUNDLE DESIGNED HELICAL BUNDLE	
Coumpound	RECEPTOR; CHAIN: A, B; ANTIBODY DESIRE-1; CHAIN: L, H;	MHC CLASS I HLA-4; CHAIN: A; BETA-2 MICROGLOBULIN; CHAIN: B; TAX PEPTIDE P6A; CHAIN: C; HMAN T-CELL RECEPTOR; CHAIN: D; HLA-A 0201: CHAIN: E:	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	ALPHA, BETA T-CELL RECEPTOR CHAIN: A, B;	LIGASE(SYNTHETASE) SERYL-TRNA SYNTHETASE (E.C.6.1.1.11) (SERINE-TRNA LIGASE) 1SES 3 COMPLEXED WITH SERYL-HYDROXAMATE- AMP 1SES 4	DHPI; CHAIN: NULL;	
SeqFold Score				51.61			
PMF Score		0.72	0.98		0.06	0.01	
Verify Score		0.34	0.37		0.01	0.56	
PSI BLAST Score		3.4e-31	1.2e-32	1.2e-32	0.0027	5.1e-05	
End AA		136	127	146	138	207	
Start AA		23	21	21	70	164	
Chain ID		Ω .	A	А	A		
PDB ID		lqm	Itcr	Itcr	lses	4hb1	
SEQ ID NO:		629	619	679	089	089	

100000

TABLE 6

SEQ ID	Position of The Last	Maximum Score	Mean Score
NO:	Amino Acid of The Signal		
342	1-13	0.981	0.764
343	1-46	0.978	0.754
344	1-34	0.954	0.756
345	1-45	0.981	0.652
346	1-22	0.982	0.882
347	1-13	0.981	0.764
348	1-27	0.992	0.969
349	1-15	0.909	0.589
350	1-33	0.961	0.864
351	1-17	0.974	0.943
353	1-20	0.957	0.874
354	1-20	0.972	0.771
355	1-28	0.941	0.755
356	1-22	0.932	0.802
357	1-20	0.895	0.595
358	1-17	0.884	0.588
359	1-16	0.988	0.881
360	1-26	0.937	0.784
361	1-29	0.981	0.864
362	1-26	0.968	0.806
363	1-22	0.968	0.806
364	1-29	0.956	0.765
365	1-21	0.992	0.929
370	1-46	0.978	0.754
380	1-34	0.954	0.756
391	1-31	0.960	0.773
399	1-45	0.981	0.652
408	1-22	0.982	0.882
409	1-42	0.993	0.715
411	1-30	0.966	0.767
423	1-18	0.997	0.971
430	1-13	0.981	0.764
435	1-45	0.890	0.631
438	1-27	0.992	0.969
466	1-33	0.961	0.864
472	1-45	0.987	0.658
473	1-20	0.992	0.967
502	1-20	0.957	0.874
503	1-21	0.989	0.945
506	1-42	0.980	0.577
511	1-20	0.972	0.771
516	1-28	0.941	0.755
517	1-28	0.941	0.755
518	1-12	0.907	0.779
522	1-21	0.958	0.779
527	1-15	0.970	0.875
538	1-20	0.895	0.595
542	1-31	0.987	0.895
545	1-30	0.971	0.889
552	1-17	0.884	0.588
562	1-23	0.965	0.817
564	1-29	0.933	0.725
575	1-28	0.972	0.870

SEQ ID	Position of The Last	Maximum Score	Mean Score
NO:	Amino Acid of The Signal		
577	1-17	0.966	0.905
586	1-26	0.921	0.587
595	1-20	0.938	0.631
606	1-18	0.901	0.763
611	1-20	0.940	0.693
615	1-26	0.937	0.784
617	1-22	0.972	0.745
618	1-15	0.930	0.748
619	1-35	0.906	0.600
622	1-29	0.981	0.864
629	1-19	0.976	0.916
630	1-27	0.973	0.931
631	1-29	0.950	0.629
632	1-19	0.969	0.913
633	1-21	0.956	0.823
637	1-17	0.976	0.938
640	1-18	0.991	0.978
645	1-26	0.968	0.806
646	1-20	0.972	0.828
647	1-27	0.893	0.567
648	1-21	0.994	0.959
649	1-20	0.945	0.891
650	1-21	0.984	0.858
651	1-27	0.891	0.593
654	1-40	0.955	0.703
668	1-22	0.968	0.806
671	1-23	0.982	0.945
672	1-23	0.982	0.945
675	1-32	0.955	0.617
676	1-23	0.936	0.677
679	1-20	0.937	0.859
680	1-29	0.956	0.765
681	1-23	0.968	0.819

TABLE 7

SEQ ID NO:	Chromosomal Location
SEQ ID NO:	17
2	10
3	11
4	4
5	15q25
6	3
7	3
9	
11	12
12	
13	17pter-p13.1
14	
15	16p13.3
16	1 12-12
17	12p13
20	21q22.3
	14
21 22	7q22
22 23	9
23	5q31 8p23-p22
25	8p23-p22
25 26	11 X
20 27	X
28	X
28 29	15q14
30	10q24
30	17q21
31 32	11
33	8
33	5q34
35	6 .
33	10
40	8q24
40	4q13.3
41	10
44	20q11.22-q12 12
47	12
48	4
49	19
50	19
51	4
52	17
55	14
56	1
57	11
57	17p13.3 5p14.2-q31.3
58 59	5p14.2-q31.3
39	7q11.2 15
60	15
61	19q13.3
62	6 5
63	5
64	7
65	22 12q24.3
66	12q24.3

SEQ ID NO:	Chromosomal Location
69	15
70	22q13.2
71	16
72	7g31.1
75	10
76	18
77	15
78	18q
79	6q14
80	11p15
81	5p13.3-q21.3
83	7q33
84	1932
85	14
87	11q12-q13.1
89 90	22
	1
91 92	1p36.13
92	7p14
93	10cen-q26.11
95	19
96	17
97	22q11.2 6p22.3
98	
99	3 8
100	11
101	2
102	7p13-p11.2
103	15q21-q22
104	15
105	9q22.1-q22.3
106	Xq13.1
107	20
108	5
109	5
· 110	16q23
111	1p32-p35
112	9
113	Xq22
114	15
115	8q22-q23
117	6p21.3
118	16p13.3
119	15
120	16
121	2q37
123	8q22-q23
124	19q13.1
126	20p12.2-p11.22
127	8
128	12pter-p13.31
129	12pter-p13.31
	18p11.22-p11.21
133 134	1q32.3-q41
134	19q13.4

SEQ ID NO:	Chromosomal Location
135	16
136	17
137	17pter-p13.1
139	7
140	8
141	Xp11.4-p11.21
142	1
143	6
144	5p14-15
145	14
146	14
147	20
148	22
149	19
150	17
151	15
152	15
154	6
155	10
156	12pter-p13.31
160	5p15.2
161 162	14q11.2
163	7q35
164	15
166	12
168	6q 18
169	7
170	7
171	6p12.1-21.1
172	6p12.1-21.1
173	15q22.1-q22.31
175	22q13.1
176	22q13.1
177	22q13.2-q13.31
178	11cen-q12.1
179	5
180	11
184	17q21.3
185	11
188	20
189	10
190	4p16
191	4
192 193	4
193	12
194	9
196	17p11.2
197	6
198	5
199	17
200 202	6q16.1-q16.3
202	1
203	2q13
203	19
407	19

SEQ ID NO:	Chromosomal Location
211	19
212	q25-26
216	19q13.3
217	21q11.2
218	. Xq21.3-q22
219	6
221	14q11.2
222	5q32
224	13
225	3q13.3-q21
226	6q23-q24
227	17
228	17
231	14
232	22
233	19
234	5q11.2
237	7q22
241	19
242	15
244	1p22
246 248	3p21.1-9
248	p12.2-13
249	10
250	19p13.3
253	19p13.3
255	4
259	9
260	5q31
262	8
264	1q32.1-q41
267	10
269	11
272	5q34
274	19
275	3
279	17
280	2
286	22q13.1
287	7
288	19q13.3-q13.4
291	2p12
292	14
293	14q31
294	11p15.5
296	7p14-p13
298	7q35-q36
299	20
300	9
302	7q22
305	14q11.2
306	11
307	14q11.2
308	14q11.2
309	7q35

SEQ ID NO:	Chromosomal Location
313	p34.3-36.11
315	17 :
316	15
317	12
318	22q11.2
319	6pter-p22.1
322	22q
323	10
326	X
328	1
329	14q11.2
330	6p21.3
331	6p21.3
332	19q13.3
333	X
334	7q31.3-q32
337	3p21.3
338	14q11.2
339	9
341	2

TABLE 8

of Full-length Nucleotide Sequence Peptide Sequence Application USSN 09/714,936 1 342 1 2 343 4 3 344 5 4 345 7 5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 <th>SEQ ID NO:</th> <th>SEQ ID NO: of Full-length</th> <th>SEQ ID NO: in Priority</th>	SEQ ID NO:	SEQ ID NO: of Full-length	SEQ ID NO: in Priority
Nucleotide Sequence 09/714,936 1 342 1 2 343 4 3 344 5 4 345 7 5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366	of Full-length	Peptide Sequence	Application USSN
1 342 1 2 343 4 3 3444 5 4 345 7 5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367	Nucleotide		09/714,936
2 343 4 3 3444 5 4 345 7 5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370	Sequence		
3 344 5 4 345 7 5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369	1	342	1
4 345 7 5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370	2	343	4
5 346 8 6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371	3	344	5
6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 338 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 30 371 43 31 372 44 32 373 <td>4</td> <td>345</td> <td>7</td>	4	345	7
6 347 10 7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 338 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 30 371 43 31 372 44 32 373 <td>5</td> <td>346</td> <td>8</td>	5	346	8
7 348 11 8 349 12 9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 336 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 33 374 <td></td> <td></td> <td></td>			
9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 32 373 45 33 374 46 34 375 47 35 376 48 36	7	348	11
9 350 13 10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 32 373 45 33 374 46 34 375 47 35 376 48 36	8	349	
10 351 14 11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 33 374 46 34 375 47 35 376 48 36 377	9		
11 352 15 12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 32 373 45 33 374 46 34 375 47 35 376 48 36 377	10		
12 353 17 13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 32 373 45 33 374 46 34 375 47 35 376 48 36 377 49 37 378 50 38 379 51 39 <td< td=""><td>11</td><td></td><td></td></td<>	11		
13 354 18 14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 32 373 45 33 374 46 34 375 47 35 376 48 36 377 49 37 378 50 38 379 51 39 380 52 40 <td< td=""><td>12</td><td></td><td></td></td<>	12		
14 355 19 15 356 20 16 357 21 17 358 22 18 359 25 19 360 29 20 361 30 21 362 32 22 363 34 23 364 36 24 365 37 25 366 38 26 367 39 27 368 40 28 369 41 29 370 42 30 371 43 31 372 44 32 373 45 33 374 46 34 375 47 35 376 48 36 377 49 37 378 50 38 379 51 39 380 52 40 381 53 41 <td< td=""><td>13</td><td></td><td></td></td<>	13		
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48 389 61 49 390 62			
49 390 62			
50 391 63			
	50	391	63

SEQ ID NO:	SEQ ID NO: of Full-length	SEQ ID NO: in Priority
of Full-length	Peptide Sequence	Application USSN
Nucleotide		09/714,936
Sequence		
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52	393	65
53	394	66
54	395	67
55	396	68
56	397	69
57	398	70
58	399	71
59	400	72
60	401	73
61	402	74
	403	75
63	404	76
64	703	. 77
66	406	
67		79
68	408	80
69	410	81
70	411	82
71	412	83 84
72	413	85
73	414	86
74	415	87
75	416	88
76	417	89
77	418	90
78	419	91
79	420	92
80	421	93
81	422	94
82	423	95
83	424	96
84	425	97
85	426	98
86	427	99
87	428	100
88	429	101
89	430	102
90	431	103
91	432	104
92	433	105
93	434	106
94	435	107
95	436	108
96	437	109
97	438	110
98	439	111
99	440	112
100	441	113
101	442	114
102	443	115
103	444	116

SEQ ID NO:	CEO ID NO. of E. II I	CEC VE NO : TO : '
	SEQ ID NO: of Full-length	SEQ ID NO: in Priority
of Full-length	Peptide Sequence	Application USSN
Nucleotide		09/714,936
Sequence		
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106	447	119
107	448	120
108	449 ·	121
109	450	122
110	451	123
111	452	124
112	453	125
113	454	126
114	455	127
115	456	
116	457	128
117		129
	458	130
118	459	131
119	460	132
120	461	133
121	462	134
122	463	135
123	464	136
124	465	137
125	466	138
126	467	139
127	468	140
128	469	141
129	470	142
130	471	143
131	472	144
132	473	145
133	474	146
134	475	
135	476	147
136		148
	477	149
137	478	150
138	479	151
139	480	152
140	481	153
141	482	154
142	483	155
143	484	156
144	485	157
145	486	158
146	487	159
147	488	160
148	489	162
149	490	163
150	491	164
151	492	165
152	493	166
153	494	
154	494	167
155		168
156	496	169
130	497	170

SEQ ID NO: of Full-length Nucleotide Sequence	SEQ ID NO: of Full-length Peptide Sequence	SEQ ID NO: in Priority Application USSN 09/714,936
157	498	171
158	499	172
159	500	173
160	501	174
161	502	175
162	503	176
163	504	177
164	505	178
165	506	179
166	507	180
167	508	181
168	509	. 182
169	510	183
170	511	184
171	512	185
172	513	186
173	514	187
174	515	188
175	516	189
176	517	190
177	518	191
178	519	192
180	520	193
181	521	194
182	522	195
183	523 524	196
184	525	197
185	526	198
186	527	199
187	528	200
188	529	201 202
189	530	203
190	531	203
191	532	204
192	533	206
193	534	207
194	. 535	208
195	536	209
196	537	210
197	538	211
198	539	212
199	540	213
200	541	214
201	542	215
202	543	216
203	544	217
204	545	218
205	546	219
206	547	220
207 ·	548	221
208	549	222
209	550	223

SEQ ID NO:	SEQ ID NO: of Full-length	SEQ ID NO: in Priority
of Full-length	Peptide Sequence	Application USSN
Nucleotide		09/714,936
Sequence		
210	551	224
211	552	225
212	553	226
213	554	227
214	555	228
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216	557	230
217	558	231
218	559	232
219	560	233
220 221	561	234
222	562	235
223	563 564	236
224		237
225	565 566	238
226		239
227	567 568	240
228	569	241
229	570	242 244
230	571	244
231	572	246
232	573	247
233	574	248
234	575	249
235	576	250
236	577	251
237	578	252
238	579	253
239	580	254
240	581	255
241	582	256
242	583	257
243	584	258
244	585	259
245	586	260
246	587	261
247	588	262
248	589	263
249	590	265
250	591	266
251	592	267
252	593	268
253	594	269
254	595	270
255	596	272
256	597	273
257	598	275
258	599	276
259	600	277
260	601	278
261	602	279
262	603	280

SEQ ID NO: of Full-length Nucleotide Sequence	SEQ ID NO: of Full-length Peptide Sequence	SEQ ID NO: in Priority Application USSN 09/714,936
263	604	281
264	605	282
265	606	283
266	607	+ **
	608	284
267 268	609	285
269	610	286
270	611	287
271	612	288
272	613	290
273	614	291
274		292
275	615	293
276	616	294
277	617	295
278	618	296 .
279	619	297
	620	298
280 281	621	299
282		300
	623	301
283	624	302
285	625	303
	626	304
286 287	627	305
288	628	306
289	629	307
290	630	308
290	631	309
292	632	310
293	633	311
294	634	312
295	635	313
296	636	314
297	637	315
298	638	316
298	639	318
300	640	319
301	641	320
302	642	321
	643	322
303	644	323
304	645	324
305 306	646	325
307	647	326
	648	327
308 309	649	328
	650	329
310	651	330
311	652	331
312	653	332
313	654	333
314	655	334
315	656	335

SEQ ID NO: of Full-length Nucleotide Sequence	SEQ ID NO: of Full-length Peptide Sequence	SEQ ID NO: in Priority Application USSN 09/714,936
316	657	336
317	658	337
318	659	338
319	660	339
320	661	340
321	662	341
322	663	342
323 `	664	343
324	665	344
325	666	345
326	667	346
327	668	347
328	669	348
329	670	349
330	671	351
331	672	352
332	673	353
333	674	354
334	675	355
335	. 676	356
336	677	357
337	678	358
338	679	359
339	680	360
340	681	361
341	682	362

WHAT IS CLAIMED IS:

An isolated polynucleotide comprising a nucleotide sequence selected from the group 1. consisting of SEQ ID NO: 1-341, a mature protein coding portion of SEQ ID NO: 1-341, an active domain coding portion of SEQ ID NO: 1-341, and complementary sequences thereof.

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2. An isolated polynucleotide encoding a polypeptide with biological activity, wherein said polynucleotide hybridizes to the polynucleotide of claim 1 under stringent hybridization conditions.

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- An isolated polynucleotide encoding a polypeptide with biological activity, wherein said polynucleotide has greater than about 90% sequence identity with the polynucleotide of claim 1.
 - 4. The polynucleotide of claim 1 wherein said polynucleotide is DNA.

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- 5. An isolated polynucleotide of claim 1 wherein said polynucleotide comprises the complementary sequences.
- 6. A vector comprising the polynucleotide of claim 1.

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- 7. An expression vector comprising the polynucleotide of claim 1.
- A host cell genetically engineered to comprise the polynucleotide of claim 1. 8.
- 25 9. A host cell genetically engineered to comprise the polynucleotide of claim 1 operatively associated with a regulatory sequence that modulates expression of the polynucleotide in the host cell.
- 10. An isolated polypeptide, wherein the polypeptide is selected from the group consisting 30 of:
 - (a) a polypeptide encoded by any one of the polynucleotides of claim 1;
 - (b) a polypeptide encoded by a polynucleotide hybridizing under stringent conditions with any one of SEQ ID NO: 1-341; and

- (c) a polypeptide of any one of SEQ ID NO: 342-682.
- 11. A composition comprising the polypeptide of claim 10 and a carrier.
- 5 12. An antibody directed against the polypeptide of claim 10.

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- 13. A method for detecting the polynucleotide of claim 1 in a sample, comprising:
- a) contacting the sample with a compound that binds to and forms a complex with the polynucleotide of claim 1 for a period sufficient to form the complex; and
- b) detecting the complex, so that if a complex is detected, the polynucleotide of claim 1 is detected.
- 14. A method for detecting the polynucleotide of claim 1 in a sample, comprising:
- a) contacting the sample under stringent hybridization conditions with nucleic acid primers that anneal to the polynucleotide of claim 1 under such conditions;
 - b) amplifying a product comprising at least a portion of the polynucleotide of claim 1; and
- detecting said product and thereby the polynucleotide of claim 1 in the 20 sample.
 - 15. The method of claim 14, wherein the polynucleotide is an RNA molecule and the method further comprises reverse transcribing an annealed RNA molecule into a cDNA polynucleotide.

16. A method for detecting the polypeptide of claim 10 in a sample, comprising:

- a) contacting the sample with a compound that binds to and forms a complex with the polypeptide under conditions and for a period sufficient to form the complex; and
- b) detecting formation of the complex, so that if a complex formation is detected, the polypeptide of claim 10 is detected.

23. The collection of claim 22, wherein the collection is provided on a nucleic acid array.

- 24. The collection of claim 23, wherein the array detects full-matches to any one of the polynucleotides in the collection.
 - 25. The collection of claim 23, wherein the array detects mismatches to any one of the polynucleotides in the collection.
- 10 26. The collection of claim 22, wherein the collection is provided in a computer-readable format.
 - 27. A method of treatment comprising administering to a mammalian subject in need thereof a therapeutic amount of a composition comprising a polypeptide of claim 10 or 20 and a pharmaceutically acceptable carrier.
 - 28. A method of treatment comprising administering to a mammalian subject in need thereof a therapeutic amount of a composition comprising an antibody that specifically binds to a polypeptide of claim 10 or 20 and a pharmaceutically acceptable carrier.

15